

GRAS Notice (GRN) No. 640

<http://www.fda.gov/Food/IngredientsPackagingLabeling/GRAS/NoticeInventory/default.htm>

**ORIGINAL SUBMISSION**

# 640



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March 8, 2016

Paulette Gaynor, Ph.D.  
Office of Food Additive Safety (HFS-200)  
Center for Food Safety and Applied Nutrition  
Food And Drug Administration  
5100 Paint Branch Parkway  
College Park, MD 20740-3835

GRN 000640

Dear Dr. Gaynor:

In accordance with 21 CFR 170.36 (62 FR 18960; April 17, 1997), Choco Finesse, LLC is hereby submitting notice of a determination based on scientific procedures that the use of H-EPG-05, as defined in the enclosed documents, is generally recognized as safe (GRAS) under specific conditions of use as an ingredient in multiple food categories, and that it is therefore exempt from the premarket approval requirement of the Federal Food, Drug, and Cosmetic Act.

Enclosed please find an electronic copy of the GRAS notice, which includes a comprehensive summary of data supporting the safety of the ingredient and the signed statement of an expert panel regarding the value of these data in supporting a GRAS determination.

My contact information is provided below. Please feel free to contact me<sup>1</sup> by phone or e-mail if you have any questions regarding this GRAS notice.

Sincerely,

(b) (6)

David Rowe  
President  
Phone: 317-694-3601  
Email: [drowe@chocofinesse.com](mailto:drowe@chocofinesse.com)

<sup>1</sup> Please note that Choco Finesse, LLC has authorized Dr. David Bechtel ([David.Becht@Intertek.com](mailto:David.Becht@Intertek.com)) from Intertek Scientific & Regulatory Consultancy, located at 100 Davidson Ave., Suite 102, Somerset, NJ 08873, to engage in discussions about any issues related to the enclosed GRAS notice. Dr. Bechtel may be reached by e-mail (shown above), by telephone at (908) 429-9202, or by FAX at (908) 429-9260.



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GRAS Exemption Claim

Choco Finesse, LLC has determined that the use of H-EPG-05 under specific conditions of use as an ingredient in multiple food categories entails a reasonable certainty of no harm and is generally recognized as safe (GRAS) based on scientific procedures. Consequently, it is exempt from the premarket approval requirement of the Federal Food, Drug, and Cosmetic Act.

(b) (6)  
[Redacted Signature]

Signature \_\_\_\_\_

Date 8 March 2016

**David Rowe**  
**President**

Name and Address of Notifier

Choco Finesse, LLC  
5019 N. Meridian Street  
Indianapolis, IN 46208

Contact Name: David Rowe  
Phone: 317-694-3601  
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GRAS Substance

The subject of this GRAS notice is H-EPG-05 marketed by Choco Finesse, LLC. The product is manufactured under current good manufacturing practices (cGMP) (21 CFR, part 110). Inspections and testing are performed at various points during the manufacturing process; every

lot of H-EPG-05 is tested for compliance with the established specifications (e.g., heavy metals, solid fat content, fatty acid composition and microbiological activity).

## Intended Use and Projected Consumer Exposure

Choco Finesse, LLC intends to market H-EPG-05 for use as a fat replacer at levels up to 38% (w/w expressed on a fat basis) in spreadable and baked goods as specified in Table 1.

**Table 1 Summary of proposed food uses and maximum use levels**

Food Category	Proposed Food-Uses	Percent (%) EPG Inclusion Expressed on Fat Basis
Baked goods and baking mixes	Biscuits	4-12
	Breads, specialty, flavored	2-8
	Brownies	6-17
	Cakes	4-14
	Cookies	7-21
	Crepes	2-20
	Desserts excluding cakes, cookies, brownies	2-20
	Doughnuts	5-12
	Muffins	2-13
	Pastry	6-15
	Pastry crusts and pies	7-25
Frozen dairy desserts and mixes	Ice cream	4-12
Grain products and pastas	Granola and other snack bars (e.g., pumpkin-based) <sup>c</sup>	4-17
Gravies and sauces	Pasta sauces (cream and tomato based)	3-8
Nut and nut products	Nut butters and nut spreads	10-38
Soft candy	Candy bars	4-17

## Basis for GRAS Determination

To make the GRAS determination, Choco Finesse, LLC compiled information about the substance, specifications, manufacturing, proposed uses, and evidence of safety into a comprehensive dossier (GRAS Dossier); and sought the opinion of qualified experts (*i.e.*, expert panel) in determining whether there is consensus among their peers that the use of this substance as described entails a reasonable certainty of no harm and is generally recognized as safe based on the available scientific evidence.

All data and information that are the basis for this GRAS determination are available for FDA's review and copying at reasonable times at Intertek Scientific & Regulatory Consultancy, 100 Davidson Ave., Suite 102, Somerset, NJ 08873, and will be sent to FDA upon request.



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## GRAS NOTICE

SUMMARY OF DATA SUPPORTING A DETERMINATION  
THAT THE USE OF ESTERIFIED PROPOXYLATED GLYCEROL (EPG)  
IN SPREADABLE AND BAKED GOODS IS  
GENERALLY RECOGNIZED AS SAFE (GRAS)

**Submitted to:**

Food and Drug Administration  
Center for Food Safety and Applied Nutrition  
Office of Food Additive Safety

**By:**

Choco Finesse, LLC  
5019 N. Meridian Street  
Indianapolis, Indiana  
46208

March 8, 2016

## **GRAS EXPERT PANEL STATEMENT**

**SUMMARY OF DATA SUPPORTING A DETERMINATION  
THAT THE USE OF ESTERIFIED PROPOXYLATED  
GLYCEROL (EPG) IN SPREADABLE AND BAKED GOODS IS  
GENERALLY RECOGNIZED AS SAFE (GRAS)**

## Expert Panel Opinion Statement Concerning the Generally Recognized as Safe (GRAS) Status of the Use of Esterified Propoxylated Glycerol (EPG) in Spreadable and Baked Goods

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Choco Finesse, LLC (hereafter referred to as Choco Finesse) has commissioned an independent panel of recognized experts (hereafter referred to as the Expert Panel) to issue an opinion whether the use of H-EPG-05 as a fat replacer at levels up to 38% (w/w expressed on a fat basis), in spreadable and baked goods is GRAS. To facilitate the review, each Expert Panel member received a comprehensive package describing the intended use, manufacturing, specifications, analytical data, exposure estimates and data supporting the safety of EPGs.

In evaluating the GRAS status of H-EPG-05, the Expert Panel considered that:

- Choco Finesse previously submitted a notification of GRAS status for the use of H-EPG-05 as a fat replacer at levels up to 34.5% (w/w) in confectionary applications to the United States Food and Drug Administration (FDA); the FDA reviewed this GRAS notification (GRN No. 583), with no resulting questions.
- The version of H-EPG-05 that is the subject of the current GRAS evaluation differs from the version of H-EPG-05 for confectionary applications (GRN No. 583), in that one-fourth to one-third of the saturated fatty acids have been replaced with unsaturated fatty acids. Although substitution with unsaturated fatty acids does not alter the digestibility of H-EPG-05, it imparts different functional properties (*i.e.*, softer and spreadable). Due to these new properties, H-EPG-05 is now suitable for use in spreadable and baked goods (*i.e.*, baked goods and baking mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, and soft candy). Therefore the proposed uses differ from those identified in GRAS 583.
- Based on the currently considered intended use levels and representative food categories from the 2009-2010 National Health and Nutrition Examination Survey (NHANES), male adults would be expected to have the highest 90<sup>th</sup> percentile all-user intake, 9.2 g EPG/person/day, on an absolute basis.
- Analytical data for EPG-05 show that the final EPG product is consistently manufactured to food-grade specifications.
- Unlike olestra, EPG is not strongly hydrophobic and exhibits far less interaction with fat-soluble substances including fat soluble vitamins. Consistent with this, the Experts noted that vitamin fortification of animal diets was not required in any of the EPG preclinical safety studies including lifetime studies in rats and mice as well as up to 3 generations in reproductive and development studies. This differed from studies conducted with olestra, which required vitamin fortification.

## **Expert Panel Opinion Statement Concerning the Generally Recognized as Safe (GRAS) Status of the Use of Esterified Propoxylated Glycerol (EPG) in Spreadable and Baked Goods**

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Use of H-EPG-05 in spreads, cooking and baking requires that it be as resistant to oxidation and thermal decomposition as current edible fats, oils and shortenings. The Panel acknowledged that results of an Oxidative Stability Index (OSI) test demonstrated that the EPG samples for the proposed uses exhibit superior resistance to oxidation compared to shortenings (*i.e.*, Cisco<sup>®</sup> shortenings and Clear Valley<sup>®</sup> oils) commonly used in cooking and baking.

During its previous GRAS evaluation of H-EPG-05 for confectionary applications, the Expert Panel had noted that EPGs have been studied extensively, and reviewed a number of both published and unpublished safety studies.

Ingested EPG is not absorbed or metabolized to any significant degree. The results of rat studies with radiolabeled EPG showed poor absorption from the gastrointestinal tract. About 70-80% of an oral dose was recovered in the feces, 10-20% in the urine, and the remaining fraction was recovered in expired air, presumably as carbon dioxide; no accumulation in tissues has been observed.

As a substance that passes through the gastrointestinal tract largely unchanged, ingested EPG was not expected to result in systemic toxicity. Indeed, the Expert Panel acknowledged that experimental animal studies with EPG showed no acute (single-dose), subchronic (90-day), or chronic (1-year) oral toxicity; genotoxicity; carcinogenicity (2-year); skin or eye irritation; dermal sensitization; or adverse effects on reproduction and offspring development. Animals in these studies were exposed to diets containing a constant level of 5% EPG (w/w) or adjusted to deliver up to 5 g/kg of body weight/day. However, additional consideration was given to possible undesirable effects resulting from its presence in the intestine; specifically, possible interference with the absorption of nutrients (*e.g.*, lipid-soluble vitamins) and passive oil leakage in stools, two effects previously associated with olestra consumption. Among the factors the Expert Panel considered was the moderate organic nature and solubility of EPG. EPG is not severely hydrophobic, exhibiting an octanol/water partition coefficient ( $K_{ow}$ ) value in the range of 3.2-3.4, similar to the triglyceride fats it is intended to replace. This is in striking contrast to olestra, which has a  $K_{ow}$  in excess of 40.

The Expert Panel also previously recognized that administration of high concentrations of EPG in the diet was associated with some effects on fat-soluble nutrients in experimental animals and humans, an effect possibly related to the presence of unabsorbed material in the gastrointestinal tract, with EPG acting as a lipid "sink" or additional "compartment" for the distribution of lipid-soluble substances. In evaluating the significance of these findings, the Expert Panel considered it reassuring that: (1) none of the animals in any of the studies exhibited clinical signs or microscopic evidence of vitamin deficiency, and there was no evidence that the vitamin levels were significantly



## **Expert Panel Opinion Statement Concerning the Generally Recognized as Safe (GRAS) Status of the Use of Esterified Propoxylated Glycerol (EPG) in Spreadable and Baked Goods**

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different from those reported in the published literature for normal control animals. In long-term EPG studies, the blood and tissue levels of fat-soluble vitamins increased and stabilized over time to fall within normal ranges reported in the literature; (2) the association between EPG consumption and effects on some fat-soluble nutrients seen in an 8-week dietary study in humans was uncertain. Beta-carotene was lower, for example, but showed no apparent relationship to EPG concentration (more evident at 10 and 40 g/day than at 25 g/day), and 25-OH D<sub>3</sub> (vitamin D, ergocalciferol) rose unexpectedly, but only in the control (margarine only) group. The Expert Panel concluded that the projected consumer exposure (90<sup>th</sup> percentile of 19.96 g/day or less) from the anticipated initial food uses would not be expected to significantly affect normal absorption of lipid-soluble nutrients. Similarly, oily stools and other gastrointestinal symptoms occasionally seen at the highest EPG dietary concentrations (up to 150 g/day) were not considered by the Expert Panel to be likely to occur at the much lower intended intake levels nor were they viewed as toxic effects.

During its previous deliberation, the Expert Panel examined the clinical chemistry results of multiple human range-finding tolerance studies and noted that serum transaminase (aspartate aminotransferase and/or alanine aminotransferase) levels exceeded the normal range in some subjects receiving EPG in amounts between 60 and 150 g/day. Likewise, high-density lipoprotein (HDL) levels below the normal range were reported in some individuals receiving EPG in amounts between 60 and 150 g EPG/day for up to 18 days. Following a more detailed review, the Expert Panel found that the occasional moderate increase in measured serum transaminase values often occurred in a transient manner, rising briefly and then returning to normal ranges. The Panel was uncertain whether this was an adaptive response but found it reassuring that this response was not observed in more extended clinical studies designed to also measure vitamin and nutrient status. Likewise, no effect on serum transaminase activity was reported in any of the animal studies including lifetime studies in rats and mice, nor was there reported any evidence of liver damage or related pathology in these investigations. For these reasons the Expert Panel concluded that the excursion of transaminase activity observed in some preliminary clinical investigations at doses exceeding 60 g/day, was not observed at lower doses or in extended preclinical safety investigations. Therefore, the Panel concluded it was not relevant to the approximate 10 g or less intake of EPG expected from the uses under consideration.

With regard to a decline in serum HDL observed in some subjects in early studies at high doses, the Panel previously noted that the effect was small and not reported in subsequent studies at lower doses for more extended periods. Likewise, the Panel noted that studies in micropigs, a species considered to be a good model for human digestion, for up to 1 year did not produce evidence of an effect on serum HDL. The Panel also noted reports that when humans are placed on a low fat diet they can exhibit a transient drop in serum HDL values and concluded that this effect may account

## **Expert Panel Opinion Statement Concerning the Generally Recognized as Safe (GRAS) Status of the Use of Esterified Propoxylated Glycerol (EPG) in Spreadable and Baked Goods**

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for the observed serum HDL decline in some subjects placed on very high EPG diets which greatly reduced the available fat in their diet. Based on these considerations, the Panel concluded that the occasionally lower HDL values observed in early clinical studies was not evidence of a toxic effect and was not relevant to the intake of EPG expected from the uses under consideration.

# Expert Panel Opinion Statement Concerning the Generally Recognized as Safe (GRAS) Status of the Use of Esterified Propoxylated Glycerol (EPG) in Spreadable and Baked Goods

## Expert Panel Opinion

Having considered all the relevant information about EPG, the undersigned members of the Expert Panel conclude that there is reasonable certainty that no harm will result from the use of H-EPG-05 as a fat replacer at levels up to 38% (w/w expressed on a fat basis) in specified spreadable and baked goods [i.e., baked goods and baking mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, and soft candy], and that such use is considered GRAS based on scientific procedures.

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Vice President, New Jersey Office  
Bridgewater, NJ

7, March 2016

Date

## **GRAS DOSSIER**

**SUMMARY OF DATA SUPPORTING A DETERMINATION  
THAT THE USE OF ESTERIFIED PROPOXYLATED  
GLYCEROL (EPG) IN SPREADABLE AND BAKED GOODS IS  
GENERALLY RECOGNIZED AS SAFE (GRAS)**

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# SUMMARY OF DATA SUPPORTING A DETERMINATION THAT THE USE OF ESTERIFIED PROPOXYLATED GLYCEROL (EPG) IN SPREADABLE AND BAKED GOODS IS GENERALLY RECOGNIZED AS SAFE (GRAS)

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## 1.0 INTRODUCTION

### 1.1 Declaration of Intent

In the United States (U.S.), substances added to food are exempt from the definition of “food additive” and thus from the premarket approval requirements outlined in Section 201(s) of the Federal Food, Drug, and Cosmetic Act if their use is generally recognized as safe (GRAS). For substances not already on the list of GRAS substances<sup>1</sup> that are not considered harmful when added to food because of their intrinsic properties and/or because sufficient information about their safety exists, a proposed rule issued in 1997<sup>2</sup> provides a framework for self-determination of GRAS. A determination that a substance is GRAS requires both technical evidence of safety and a basis to conclude that this technical evidence of safety is generally known and accepted within the qualified scientific community. Self-determination of GRAS may be followed by notification to the Food and Drug Administration (FDA).

Esterified propoxylated glycerols (EPG or EPGs hereafter), are a family of fat- and oil-like substances that resemble triglycerides in structure and appearance, but they have been modified to prevent or limit their digestion when consumed in food. Due to the nature of the manufacturing process, a large number of versions of EPG are made available through modification of the fatty acid moieties of the triglyceride and the extent of the propoxylation of the glycerol. A core version, H-EPG-05 HR/SO 9:1, was selected as representative of the initial EPG-05 forms for commercial development.

Choco Finesse, LLC (hereafter Choco Finesse) previously submitted a notification of GRAS status for the use of H-EPG-05 as a fat replacer at levels up to 34.5% (w/w) in confectionary applications to the United States Food and Drug Administration (FDA); the FDA reviewed this GRAS notification (GRN No. 583), with no resulting questions. Both the previously notified version of H-EPG-05, and the version of H-EPG-05 that is the subject of the current GRAS notification, exhibit a melting point above body temperature. The version of H-EPG-05 that is the subject of the current GRAS evaluation differs from the version of H-EPG-05 for confectionary applications, in that one-fourth to one-third of the saturated fatty acids have been replaced with unsaturated fatty acids (refer to Table 2-1). Substitution with unsaturated fatty acids does not alter the digestibility of H-EPG-05 (refer to Section 2.3), however, it imparts different functional properties (*i.e.*, softer and spreadable). Due to these new properties, H-EPG-05 is now suitable for use in spreadable and baked goods. As such, Choco Finesse, LLC has self-affirmed through scientific procedures that the use of H-EPG-05 as a fat replacer at levels up to 38% (w/w expressed on a fat basis), in spreadable and baked goods such as, baked goods and baking

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<sup>1</sup> 21 CFR 182 – Substances generally recognized as safe; 21 CFR 184 – Direct food substances generally recognized as safe; 21 CFR 186 – Indirect food substances affirmed as generally recognized as safe.

<sup>2</sup> 62 FR 18938; April 17, 1997.

mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, and soft candy, is GRAS. These uses differ from those identified in GRAS 583.

According to the intake assessment, when considering intakes by all-users, male adults were determined to have the highest 90<sup>th</sup> percentile all-user intake of EPG on an absolute basis at 9.2 g/person/day. The safety and tolerability of this level of intake has been clearly demonstrated in preclinical and human studies (refer to Sections 5.0 and 6.0).

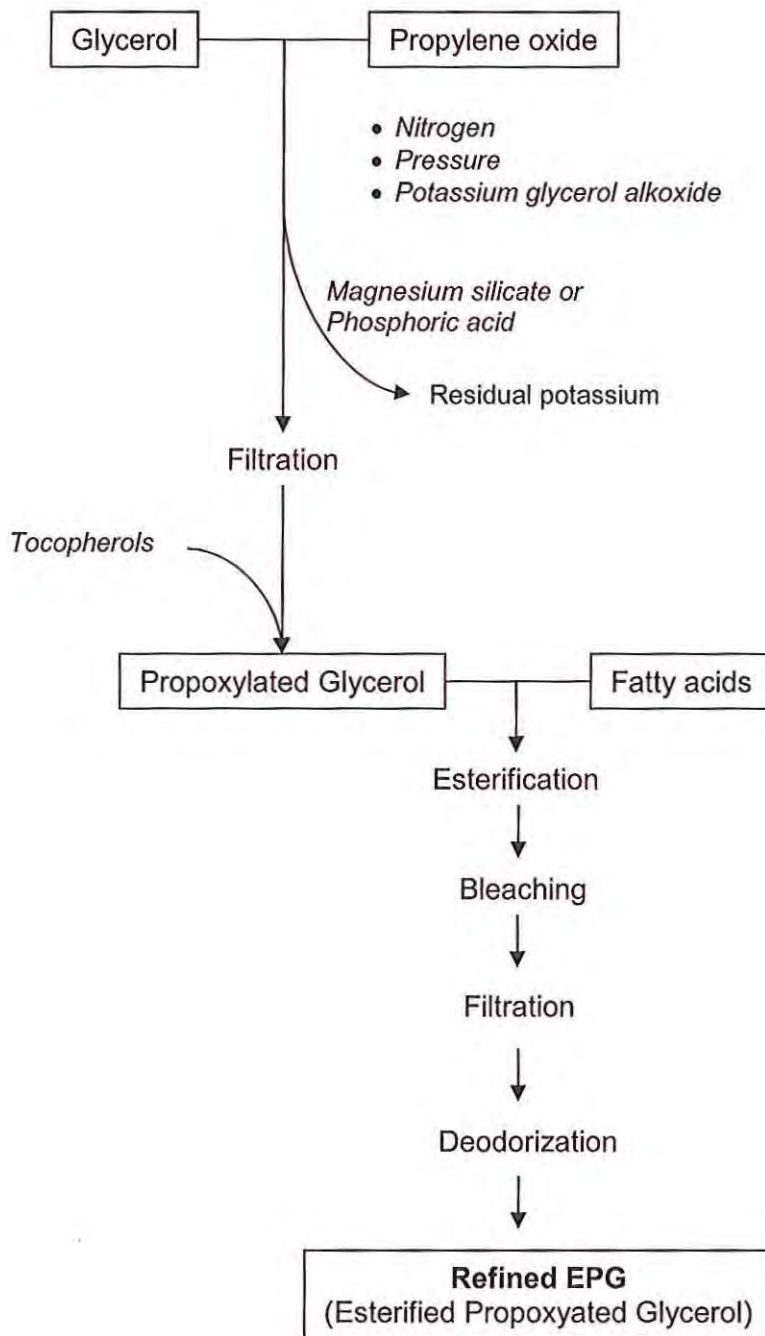
The present document summarizes the available information supporting the safety of H-EPG-05, and provided the basis for the GRAS determination.

## **2.0 MANUFACTURING AND GRAS SUBSTANCE CHARACTERIZATION**

### **2.1 Manufacturing Process**

EPG is manufactured in compliance with current Good Manufacturing Practice (cGMP) regulations. Briefly, the production of EPG consists of two basic processes: (1) propoxylation of glycerol; and (2) esterification of propoxylated glycerol with fatty acids. Propoxylation of glycerol involves reacting food grade glycerol with propylene oxide under base catalysis to form the tri-functional polyether polyol (propoxylated glycerol). The esterification is carried out without catalyst using an excess of fatty acids. The unsaturated fatty acids are derived from splitting natural edible fats and oils, while saturated fatty acids are produced by splitting fully hydrogenated edible oils. The unreacted fatty acids are removed from crude EPG by standard steam stripping under vacuum (*e.g.*, during physical refining-deodorization). The manufacturing process for EPG is also depicted in Figure 2-1.

Figure 2-1 Manufacturing process of EPG



## 2.2 Food-Grade Specifications and Chemical Analysis

The specifications for H-EPG-05 for use in spreadable and baked goods are summarized in Table 2-1.

**Table 2-1 Specifications for H-EPG-05 for use in spreadable and baked goods**

Attribute	Specification	Range	Method
Appearance	Solid, white < 30°C	Sl. off white	Visual (internal procedure)
Melting point	Mettler Dropping Point	38-43° C	AOCS Tr Ia-64
Taste	Sensory	Flavorless	Taste (internal procedure)
Texture	Sensory	Waxy	Taste (internal procedure)
Free Fatty Acid, %	% FFA as oleic	0.50 max.	AOCS Ca 5a-40
Peroxide Value	Meq.peroxide/1000g	0-1	AOCS Cd 8b-90
Anisidine Value	p-anisidine	1-10	AOCS Cd18-90
Hydroxyl Value	mg KOH/g	5.0 max.	ASTM D4274
Iodine Value	mg Iodine/g	15-30	AOCS Cd 1-25
Purity, % EPG (100-%FFA)	Esters content	>99.5	Calculation (esters – FFA)
Oxidative Stability Index (OSI), hrs.		>50	
Trace Metals, ppm	Iron	<0.1	AOCS Ca17-1
	Copper	<0.01	
	Calcium	<0.5	
	Magnesium	<0.5	
Tocopherols, ppm	Alpha	120-230	AOCS Ce 8-89
	Beta	5-40	
	Gamma	500-800	
	Delta	150-250	
	Total	900-1300	
Fatty Acid Composition	Palmitic, C16:0	0-1.0	AOCS Ce 1-62
	Stearic, C18:0	4-10	
	Oleic, C18:1	20-30	
	Linoleic, C18:2	1.0-3.0	
	Linolenic, C18:3	0-0.5	
	Arachidic, C20:0	38-48	
	Behenic, C22:0	20-30	
	Arachidic + Behenic	58-78	
	Total saturates	67-79	
Total unsaturates	21-33		
Trans fat, %	Total "trans"	<1.0*	AOCS Ce 1h-05
Solid Fat Content	@ 10°C	65-75	Cd16b-IUPAC
	@ 20°C	58-68	
	@ 25°C	43-55	
	@ 30°C	30-40	
	@ 35°C	17-25	
	@ 40°C	0-0.5	
Smoke Point	Open Cup, °C	>220	AOCS Cc 9a-48
Flash Point	Open Cup, °C	>265	

\* Trans fat contributed by Pamolyn oleic acid used for esterification of EPG. Less than 8% of the amount present is bioavailable which amounts to less than 0.05% trans fatty acid that may actually be absorbed from EPG. Conjugated linoleic acid (CLA) not included in total trans values.

**Table 2-1 Specifications for H-EPG-05 for use in spreadable and baked goods (cont'd)**

Heavy Metals	Specification	Range	Method
Arsenic		<0.05	ICP-MS/AOAC 993.14
Lead		<0.05	
Microbiological Screening	Specification	Range	Method
Aerobic Plate Count	Petrifilm	<10/g	AOAC 990.12
Coliform	Petrifilm	<10/g	AOAC 991.14
<i>E. coli</i>	Petrifilm	<10/g	
<i>Salmonella</i>	ELFA	Negative/25g	AOAC 2004.03
Yeast		<10/g	FDA-BAM, 7 <sup>th</sup> ed.
Mold		<10/g	

Analyses of non-consecutive lots of H-EPG-05 for use in spreadable and baked goods are presented in Table 2-2.

**Table 2-2 Batch analysis data for H-EPG-05 for use in spreadable and baked goods**

Attribute	Specification	Lot Numbers		
		Batch B160 7/15/15	Batch B168 8/12/15	Batch 169 8/12/15
Appearance	Solid, white < 30 °C	White, solid	White, solid	White, solid
Mettler Dropping Point, °C	38-43	38.9	38.5	38.6
Taste	Flavorless	Flavorless	Flavorless	Flavorless
Texture	Waxy	Waxy	Waxy	Waxy
Free Fatty Acid, % (as oleic)	0.50 max.	0.32	0.50	0.49
Peroxide Value	0-1	0.6	0.1	0.2
Anisidine Value	1-10	1.6	1.4	1.0
Hydroxyl Value	5.0 max.	5.0	2.7	2.7
Iodine Value	15-30	19.5	21.4	20.4
Purity, % EPG (100-%FFA)	>99.5	99.68	99.50	99.51
Trace Metals				
Iron, ppm	<0.1	<0.1	<0.1	<0.1
Copper, ppm	<0.01	<0.01	<0.01	<0.01
Calcium, ppm	<0.5	<0.5	<0.5	<0.5
Magnesium, ppm	<0.5	<0.5	<0.5	<0.5
Tocopherols				
Alpha, ppm	120-230	168	166	158
Beta, ppm	5-40	19	19	17
Gamma, ppm	500-800	691	713	598
Delta, ppm	150-250	234	246	203
Total, ppm	900-1300	1112	1114	976
Fatty Acid Composition, %				
Palmitic, C16:0	0-1.0	0.4	0.4	0.2
Stearic, C18:0	4-10	5.6	5.4	5.5
Oleic, C18:1	20-30	23.5	24.9	24.7
Linoleic, C18:2	1-3.0	1.3	1.4	1.3
Linolenic, C18:3	0-0.5	0.3	0.2	0.2
Arachidic, C20:0	38-48	44.0	43.1	43.3
Behenic, C22:0	20-30	23.4	22.9	23.0
Arachidic + Behenic	58-78	67.4	66.0	66.3
Total saturates	67-79	73.4	71.8	72.0
Total unsaturates	21-33	26.6	28.2	28.0
Trans fat, %	<1.0	0.8	0.9	0.8

**Table 2-2 Batch analysis data for H-EPG-05 for use in spreadable and baked goods (cont'd)**

Attribute	Specification	Lot Numbers		
		Batch 88b 6/11/14	Batch 89 6/11/14	Batch 91 6/19/14
Solid Fat Content				
@10°C	65-75	70.4	68.4	68.8
@20°C	58-68	61.7	59.5	60.0
@25°C	43-55	49.2	45.3	46.1
@30°C	30-40	33.4	30.4	30.7
@35°C	17-25	20.2	17.6	18.2
@40°C	0-0.5	0.13	0.04	0.16
Smoke Point, °C	>200	220	220	226
Flash Point, °C	>265	310	310	320
<b>Heavy Metals</b>				
Arsenic	<0.05	<0.006	<0.005	<0.007
Lead	<0.05	<0.006	<0.004	<0.007
<b>Microbiological Screening</b>				
Aerobic Plate Count	<10/g	<10/g	<10/g	<10/g
Coliform	<10/g	<10/g	<10/g	<10/g
<i>E. coli</i>	<10/g	<10/g	<10/g	<10/g
<i>Salmonella</i>	Negative/ 25 g	Negative/ 25 g	Negative/ 25 g	Negative/ 25 g
Yeast	<10/g	<10/g	<10/g	<10/g
Mold	<10/g	<10/g	<10/g	<10/g

### 2.3 Acid and Lipase Resistance and Caloric Availability

As detailed in the previous GRAS Notification for EPG for confectionary applications (GRN No. 583), EPG-05 versions are resistant to hydrochloric acid and pancreatic lipase, and yield 0.7 kcal/g.

### 2.4 Thermal Stability

Lower solid fat content ensures softer, less brittle texture, and the broader melting profile desired in spreads (e.g., nut butters, margarines and cooking/baking shortenings). These properties are achieved by incorporating unsaturated fatty acids up to 30% of the total fatty acid content in EPG.

It is also desirable that EPGs used in spreads, cooking and baking is as resistant to oxidation and thermal decomposition as current edible fats, oils and shortenings. A standard method for



the assessment of the comparative stability of edible oils is the Oxidative Stability Index (OSI) (Medallion Laboratories, undated). This method measures relative resistance of fats and oils to oxidation, as well as decomposition of oxidation products such as, hydroperoxides at a temperature of 110 °C (230 °F). Table 2-3 shows OSI results for 3 spreadable EPG samples; as controls, and for direct comparison, two Crisco shortenings commonly used in cooking and baking were included in the same OSI test. Results demonstrated that EPG samples exhibit superior resistance to oxidation compared to shortenings commonly used in cooking and baking.

**Table 2-3 OSI results for EPG and Crisco samples**

Sample	OSI hours @ 110°C (230°F)
Spreadable EPG B156	99.7
Spreadable EPG B160	110.1
Spreadable EPG B161	105.3
Crisco Vegetable shortening	16.9
Crisco Butter shortening	15.4

Another example of edible oils recently developed specifically for cooking and baking are high oleic varieties of canola and sunflower oils (Cargill, 2015). The published OSI data for Cargill’s Clear Valley® brand are shown in Table 2-4. The resistance of these new Cargill oils to oxidation, and the thermal degradation of oxidation compounds, is similar to Crisco shortenings and inferior to the stability of spreadable EPGs.

**Table 2-4 OSI results for Cargill’s Clear Valley® brand**

Sample	OSI hours @ 110°C (230°F)
Clear Valley® 65 High Oleic Canola Oil	11
Clear Valley® 80 High Oleic Canola Oil	20
Clear Valley® High Oleic Sunflower Oil	15

Two additional measures of the thermal stability of fats and oils are their Smoke and Flash points (ChartsBin, 2011; Institute of Shortening and Edible Oils, Inc., undated). Smoke Point is the temperature at which a sample heated in the presence of air shows the first signs of bluish smoke rising from the surface. Flash Point is the lowest temperature at which an oil generates sufficient vapors to form an ignitable mixture in air.

The values in the Table 2-5 represent typical temperatures for Smoke and Flash points as determined by the AOCS test method. As the values indicate, the thermal stability of spreadable EPGs is comparable to common vegetable oils used in cooking and baking. This is expected as EPG contains fatty acids derived from vegetable oils.

**Table 2-5 Smoke and flash points for EPG and common vegetable oils\***

Fat/Oil Type	Smoke Point, °C (°F)	Flash Point, °C (°F)
Spreadable EPG B160	220 (428)	310 (590)
Spreadable EPG B168	220 (428)	310 (590)
Spreadable EPG B169	226 (439)	320 (608)
Canola oil	236 (457)	326 (619)
Coconut oil	177 (351)	295 (563)
Cottonseed oil	216 (421)	319 (606)
Corn oil	236 (457)	325 (617)
Peanut oil	231 (448)	334 (633)
Palm oil	230 (446)	324 (615)
Soybean oil	241 (466)	330 (626)

\*Determined by AOCS Cc 9a-48 method, Cleveland Open Cup

In conclusion, spreadable EPGs are significantly more resistant to oxidation compared to edible fats and oils as measured by OSI, and comparable to edible fats and oils in thermal stability as measured by Smoke and Flash points.

### **3.0 PROPOSED USE AND ANTICIPATED CONSUMER INTAKE**

#### **3.1 Proposed Uses**

For the purpose of this GRAS notification, Choco Finesse's EPG, manufactured in accordance with GMP as specified in 21 CFR 110, may be used as a fat replacer at levels up to 38% (w/w expressed on a fat basis), in spreadable and baked goods in the following categories of foods as defined in *21 CFR §170.3(n)*: baked goods and baking mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, and soft candy. The individual proposed food uses and use levels, are summarized in Table 3-1. This information was used to estimate consumer intakes, which are discussed further below. The full intake assessment report is attached as Appendix 1 of the current report.

**Table 3-1 Summary of the Individual Proposed Uses and Use Levels for EPG in the United States (2011-2012 NHANES Data)**

Food Category	Proposed Food-Uses	Percent (%) Fat in Product <sup>a</sup>	Percent (%) EPG Inclusion Expressed on Fat Basis	Percent (%) EPG Inclusion in Final Product <sup>b</sup>
Baked goods and baking mixes	Biscuits	14 to 18	4-12	2.16
	Breads, specialty, flavored	7 to 11	2-8	0.88
	Brownies	22 to 30	6-17	5.10
	Cakes	15 to 20	4-14	2.80
	Cookies	22 to 30	7-21	6.30
	Crepes	8 to 28	2-20	5.60
	Desserts excluding cakes, cookies, brownies	8 to 28	2-20	5.60
	Doughnuts	20 to 23	5-12	2.76
	Muffins	9 to 19	2-13	2.47
	Pastry	21	6-15	3.15
	Pastry crusts and pies	23 to 36	7-25	9.00
Frozen dairy desserts and mixes	Ice cream	15 to 21	4-12	2.52
Grain products and pastas	Granola and other snack bars (e.g., pumpkin-based) <sup>c</sup>	14 to 24	4-17	4.08
Gravies and sauces	Pasta sauces (cream and tomato based)	11	3-8	0.88
Nut and nut products	Nut butters and nut spreads	32 to 55	10-38	20.90
Soft candy	Candy bars	14 to 24	4-17	4.08

<sup>a</sup> Values listed were provided by the client. These were compared to the mean values per category available in the 2011-2012 NHANES data and noted to be higher; therefore, the intake assessment conducted using the percentage of fat provided by Choco Finesse presented herein is considered to be a conservative estimate.

<sup>b</sup> Calculated by (Maximum percent fat in product) \* (Maximum percent EPG inclusion expressed on a fat basis)

<sup>c</sup> No pumpkin-based bars were identified in the NHANES dataset, therefore this food use has not been included in the present assessment.

To examine worst-case EPG exposure, the intake assessment considered the total daily intake of EPG from all food-uses, based on the maximum range of EPG inclusion expressed on a fat basis. Food codes representative of each proposed food-use were chosen from the NHANES 2011-2012 (CDC, 2014; USDA, 2014). Food codes were grouped in food-use categories according to Title 21, Section §170.3 of the Code of Federal Regulations (CFR, 2014a). Product-specific adjustment factors were developed based on data provided in the standard recipe file for the Continuing Survey of Food Intakes by Individuals (CSFII) 1994-1996, 1998

survey (USDA, 2000). All food codes included in the current intake assessment are listed in the full intake assessment report (Appendix 1).

### **3.2 Anticipated Consumer Exposure**

Estimates for the total daily intakes of EPG by various U.S. population groups from all proposed food-uses are provided in Tables 4-2 and 4-3. Table 4-2 summarizes the estimated total intake of EPG on a grams per person per day basis (g/person/day), and Table 4-3 presents these data on a per kilogram body weight per day basis (mg/kg bw/day). All-person intake refers to the estimated intake of EPG averaged over *all* individuals surveyed, regardless of whether they potentially consumed food products containing EPG, and therefore includes individuals with “zero” intakes (*i.e.*, those reporting no intake of foods like those for which EPG is intended during the 2 survey days). All-user intake refers to the estimated intake of EPG by individuals that reported consuming food products like those for which EPG is intended. Individuals were considered *users* if they consumed 1 or more food products containing EPG on either Day 1 or Day 2 of the survey.

A total of 82.4% of participants were identified as users (*i.e.*, consumers) of food products in which EPG is currently proposed for use (range of 70.3-91.7%). Estimates based on all-users revealed mean and 90<sup>th</sup> percentile EPG intakes of 3.6 g/person/day (63 mg/kg bw/day) and 8.0 g/person/day (146 mg/kg bw/day), respectively, for the total U.S. population. On an individual group basis, male adults had the greatest estimated mean and 90<sup>th</sup> percentile all-user intakes on an absolute basis (4.1 and 9.2 g/person/day, respectively). As might be expected, EPG intake estimates on a per kg body weight basis were greatest among younger (*i.e.*, smaller) individuals. Specifically, infants had the greatest projected intakes per body weight (168 and 358 mg/kg bw/day for the mean and 90<sup>th</sup> percentile, respectively). However, infants are not expected to be significant consumers of EPG-containing foods and would therefore be expected to have far smaller exposures to EPG (from only occasional consumption of EPG-containing foods) than the present (worst-case) intake assessment would indicate. None of the EPG-containing products would be targeted or marketed for consumption by infants; their exposure is expected to results from collateral ingestion of items being eaten by older persons.

Total population values are included for completeness. All-person mean and 90<sup>th</sup> percentile intakes of EPG from all proposed food-uses were estimated to be 2.9 g/person/day (52 mg/kg bw/day) and 7.3 g/person/day (128 mg/kg bw/day), respectively, for the total U.S. population. Within the individual population groups, male adults were determined to have the greatest estimated mean and 90<sup>th</sup> percentile intakes of EPG on an absolute basis, at 3.4 g/person/day and 8.7 g/person/day, respectively.

**Table 4-2 Summary of the Estimated Daily Intake of EPG from Proposed Food-Uses in the United States by Population Group (2011-2012 NHANES Data)**

Population Group	Age Group (Years)	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	% Users	n	Mean	90 <sup>th</sup> Percentile
Infants and Young Children	0 to 3	1.6	4.2	70.3	547	2.3	5.2
Children	4 to 11	3.2	7.1	91.7	1,198	3.5	7.4
Female Teenagers	12 to 19	2.2	5.5	77.6	429	2.8	5.7
Male Teenagers	12 to 19	3.0	7.7	78.3	419	3.9	9.0
Female Adults	20 and up	2.7	6.8	82.6	1,761	3.3	7.3
Male Adults	20 and up	3.4	8.7	82.5	1,645	4.1	9.2
Total Population	All Ages	2.9	7.3	82.4	5,999	3.6	8.0

**Table 4-3 Summary of the Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Proposed Food-Uses in the United States by Population Group (2011-2012 NHANES Data)**

Population Group	Age Group (Years)	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
Infants and Young Children	0 to 3	118	304	70.4	546	168	358
Children	4 to 11	115	277	91.7	1,198	125	280
Female Teenagers	12 to 19	39	102	77.2	419	50	120
Male Teenagers	12 to 19	49	130	78.2	416	63	153
Female Adults	20 and up	38	93	82.6	1,746	46	104
Male Adults	20 and up	40	103	82.5	1,630	49	107
Total Population	All Ages	52	128	82.4	5,955	63	146

Estimates for the mean and 90<sup>th</sup> percentile daily intakes of EPG from each individual food category are summarized in Tables A-1 to A-7 and B-1 to B-7 of Appendix 1 on a g/day and mg/kg body weight/day basis, respectively. The total U.S. population was identified as being significant consumers of cookies (26.4 to 46.8% users), ice cream (15.4 to 33.3% users) and pasta sauces (19.9 to 27.8% users).

In terms of contribution to total mean intake of EPG, cookies (contributed 19.5 to 31.2% to total mean intakes) and nut butters (contributed 16.5 to 28.3% to total mean intakes) were the 2 main sources of intake across all population groups on both an absolute and on a mg/kg body weight

basis. Pastries individually contributed  $\leq 1.0\%$  to total mean EPG intakes across all population groups (see Tables A-1 to A-1 and/or B-1 to B-7 of Appendix 1 for further details).

## 4.0 PHARMACOKINETICS

For completeness, the summary and detailed study results from GRN No. 583 are included herein.

### 4.1 Overview of EPG Absorption, Distribution, Metabolism and Excretion

The pharmacokinetics of two radiolabeled EPG versions [H-EPG-08 oleate (a semi-solid) and H-EPG-14 oleate (a liquid)] were evaluated in male and female CrI:CD<sup>®</sup>BR rats to determine the absorption, distribution, metabolism and excretion (ADME) profile of these materials. Two separate studies were conducted with H-EPG-08 oleate; one in which the material was <sup>14</sup>C-radiolabeled on the C<sub>1</sub>-carbon of the propylene glycol units (Section 4.2.1), and a second in which it was <sup>14</sup>C-radiolabeled on the carboxyl carbon of the fatty acid portion (Section 4.2.2). In the latter study, thin-layer chromatography (TLC) was used to confirm the presence <sup>14</sup>C-oleic acid in liver tissue extracts, as incorporation of the radiolabeled fatty acid into tissues was expected. Finally, one study was conducted with H-EPG-14 oleate radiolabeled on the C<sub>1</sub>-carbon of the propylene glycol units (Section 4.2.3).

In each study, five rats/sex/group were administered a single oral dose of 1.0 g/kg bw or 3.0 g/kg bw by gavage. A third dose group was given 1.0 g/kg bw of the radiolabeled version after two weeks of daily EPG administration of the non-labeled version at the same dose. In addition, to simulate the worse case scenario of complete absorption of the EPGs, ADME studies were conducted on rats receiving a dose of 35 mg/kg of each version intravenously in a liposome suspension. Expired air, feces, urine, organs, tissue samples and the carcass were monitored for radioactivity for up to one week after EPG administration when, essentially, there was complete recovery of the dose administered.

The results of these studies indicate that the two EPG versions evaluated were poorly absorbed from the GI tract and could not be found intact in any tissues after oral dosing. EPG-08 oleate was degraded approximately 20%, while EPG-14 oleate was degraded by approximately 10%. There was some evidence that possible bacterial degradation in the gastrointestinal (GI) tract was taking place, particularly in the colon. The pattern of distribution of the radiolabel observed in the body of the rats was consistent with GI absorption of fatty acids and the propylene glycol units modified glycerol, both of which were partially oxidized to carbon dioxide. A significant portion of the fatty acids absorbed were incorporated into triglycerides and stored in adipose tissue.

In the two studies where the propoxylated glycerol units were radiolabeled, small amounts of radiolabel were detected in the liver and other metabolically active tissues, indicating that a small portion of this material was assimilated into normal body constituents during the oxidation process. This absorption, disposition, metabolism and excretion pattern for EPG was considered predictable and similar to that which would be expected from normal triglycerides. When given intravenously in fine liposome emulsion, the two versions of EPGs tested were rapidly oxidized to fatty acids and glycerol containing propoxylated glycerol units. The disposition pattern was similar to that *via* the oral route, except that larger portions of the metabolites of the EPGs were deposited in the liver and lungs. The route by which the metabolites of the various versions of EPGs were excreted appeared to be governed by their molecular weights. The greater the molecular weight, the more of the metabolites of the EPGs excreted into the feces, and the less into the urine.

More detailed summaries of the ADME studies are provided in Section 4.2.

## 4.2 ADME Studies

### 4.2.1 Metabolism and Disposition of EPG-08 Oleate in Rats (COVANCE 6226-104; E-012)

The metabolism and disposition of oxypropylene -<sup>14</sup>C-EPG-08 oleate were studied in CrI:CD<sup>®</sup>BR rats. Forty-four animals (22/sex) were divided into five groups and exposed to oxypropylene-<sup>14</sup>C-EPG-08 oleate by gavage or intravenously. A preliminary group was exposed to a single dose of 1,000 mg/kg, and used to assess the potential for excretion of radioactivity in expired air. Since approximately 2% to 8% of oxypropylene -<sup>14</sup>C-EPG-08 oleate was recovered in the expired air of rats in the preliminary group, expired air was also collected from the rats in the remaining groups. The remaining four groups were administered a single intravenous dose (30 mg/kg), a single oral dose (1,000 or 3,000 mg/kg), or 1,000 mg/kg of non-radiolabeled EPG-08 oleate for 14 days followed by a single radiolabeled, oral administration on the 15<sup>th</sup> day. Mean recovery of radioactivity in this study is summarized in Table 4-1.

Excretion of oxypropylene -<sup>14</sup>C-EPG-08 oleate was measured in the feces, urine, and expired air (presumably as CO<sub>2</sub>). Oxypropylene -<sup>14</sup>C-EPG-08 oleate *via* i.v., was excreted predominantly in the urine with 49.9% and 59.9% of the test material excreted by males and females, respectively. Recovery of radioactivity in expired air averaged 29.9% and 19% for males and females, respectively. Recovery of test material in the feces was 9.09% and 9.73% for males and females, respectively. In contrast, the fecal route was the predominant route of excretion following oral administration of oxypropylene -<sup>14</sup>C-EPG-08 oleate, demonstrated by 66.1% to 84.2% of recovered test material in feces. Recovery of radiolabeled test material in urine ranged from 7.96% to 14.3%, while recovery in expired air ranged from 2.87% to 8.11%.

Rats were sacrificed 7 days after the administration of the radiolabeled dose and various tissues<sup>3</sup> were collected and analyzed for total radioactivity. Rats exposed intravenously had the highest concentrations of radioactivity in the spleen, liver, lungs, kidneys, and bone.

Approximately 6% of the administered substance was recovered from the tissues and carcasses of these animals. Rats exposed orally had the highest concentrations of radioactivity in the liver, kidneys, spleen, lungs, and stomach. However, tissues and carcasses of these rats accounted for less than 0.5% of the administered test material.

In conclusion, oxypropylene-<sup>14</sup>C-EPG-08 oleate was poorly absorbed *via* the oral route in rats. Administration of oxypropylene -<sup>14</sup>C-EPG-08 oleate *via* i.v. results in de-esterification of the material to form diester, monoester, and base polyol. The polyol may be further metabolized to products that are excreted in urine, incorporated into endogenous components, or eliminated as CO<sub>2</sub>.

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<sup>3</sup> Tissues included bone (femur), brain, fat, heart, kidneys, large intestine, liver, lungs, muscle (thigh), ovaries, small intestine, spleen, stomach, tail, testes and uterus.



**Table 4-1 Mean recovery (%) of radioactivity from <sup>14</sup>C-EPG-08 oleate in rats (5/group) after 7 days**

Route	Dosage (mg/kg bw)	Males								Females							
		Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total	Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total
Intravenous (single dose)	30	0.29	29.9	9.09	49.9	<0.01	1.85	4.38	95.4	0.79	19.0	9.73	59.9	<0.01	1.28	4.62	95.3
Oral (single dose)	1000	0.63	7.11	74.5	11.6	<0.01	0.52	0.12	94.5	1.17	3.85	78.9	11.4	<0.01	0.27	0.05	95.7
Oral (multiple dose; radiolabeled EPG on Day 15)	1000	3.21	8.11	66.1	14.3	<0.01	0.66	0.18	92.6	0.31	3.32	78.5	10.4	<0.01	0.22	0.04	92.8
Oral (single dose)	3000	1.16	7.19	74.7	11.0	<0.01	0.61	0.15	94.8	0.22	2.87	84.2	7.96	<0.01	0.17	0.04	95.4

<sup>a</sup>Includes cage wash/wipe with 1% trisodium phosphate and hexane.

<sup>b</sup>Ethoxyethanol:ethanolamine trap; includes backup.

#### 4.2.2 Metabolism and Disposition of EPG-08 [<sup>14</sup>C] – Oleate in Rats (HWI 6226-108; E-013)

The purpose of this study was to assess the bioavailability and extent of absorption, distribution, elimination, and biotransformation of EPG-08 [<sup>14</sup>C] - oleate administered orally to CrI:CD<sup>®</sup>BR rats (131 to 182 g). Forty treated animals were divided into 4 groups of 10 animals (5/sex) as follows: single intravenous (i.v.) low dose (30 mg/kg), single oral low dose (1,000 mg/kg), multiple oral low dose (1,000 mg/kg; 14 daily nonradiolabeled doses followed by a single radiolabeled dose on the 15<sup>th</sup> day), and single oral high dose group (3,000 mg/kg). All groups had urine, feces, expired carbon dioxide, and organic volatiles collected. The animals were sacrificed 7 days after the administration of the radiolabeled dose and various tissues<sup>4</sup> were collected and analyzed for total radioactivity. Mean recovery of radioactivity in this study is summarized in Table 4-2.

Most of the radioactivity was found in the feces in all oral dose groups with values ranging from 77.1% to 83.3% of the dose. The next most important route of elimination was expiration of CO<sub>2</sub>, representing 5.38% to 10.4% of the dose. The least significant route of elimination was excretion in urine with values ranging from 0.24% to 0.52% of the dose. In contrast, following i.v. administration expiration as CO<sub>2</sub> was the predominant route of excretion representing 47.4% and 56.9% of the dose for males and females, respectively. Excretion in urine was minor with 1.30% and 1.64% of the dose excreted by males and females, respectively. Similarly, feces contained 1.42% and 1.54% of the dose for males and females, respectively. The remainder of the radioactivity was not excreted by the 7-day sacrifice time. The highest percentage of the unexcreted dose was recovered in the residual carcass for both males (26.4%) and females (15.9%). Of the discrete tissues collected, the liver (7.62% of the dose in males and 9.12% of the dose in females), tail (2.23% of the dose in males and 1.57% of the dose in females), and fat (2.27% of the dose in males and 1.25% of the dose in females) contained the most radioactivity. Material balance was high for all dose groups, ranging from 89.6% to 96.5%.

With respect to concentrations of radioactivity in the carcass and tissues following oral administration, the highest percent of dose was recovered in the carcass (1.42 to 5.31%). Tissues accounted for 0.17 to 0.60% of the radioactivity recovered, with the fat, large intestine, small intestine, stomach, lungs, ovaries, and uterus containing the highest concentrations of all the tissues examined. The radioactivity in the liver and lung tissue of the oral dose groups was believed to most likely be the result of absorption of oleic acid or oleic acid fragments resulting from chemical hydrolysis or enzymatic attack of the oxypropylene linkage in the digestive tract. In order to substantiate this hypothesis, the lipid components of tissues were extracted and the remaining residues were subjected to protease digestion. The resulting extracts were analyzed

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<sup>4</sup> Tissues included bone (femur), brain, fat, heart, kidneys, large intestine, liver, lungs, muscle (thigh), ovaries, small intestine, spleen, stomach, tail, testes and uterus.

by liquid scintillation counting (LSC) and by TLC to characterize the  $^{14}\text{C}$ -components. The presence of  $^{14}\text{C}$ -oleic acid was confirmed by TLC profiling of liver tissue extracts from all dose groups. The distribution of radioactivity in tissue components appeared to be due to the incorporation of oleic acid. With respect to the i.v. dose, the tissues that contained the highest concentrations of radioactivity were fat, spleen, liver, lungs, and carcass.

**Table 4-2 Mean recovery (%) of radioactivity from EPG-08 <sup>14</sup>C-oleate in rats (5/group) after 7 days**

Route	Dosage (mg/kg bw)	Males								Females							
		Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total	Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total
Intravenous (single dose)	30	<0.01	47.4	1.42	1.30	0.03	26.4	14.5	91.1	0.03	56.9	1.54	1.64	0.05	15.9	14.9	90.9
Oral (single dose)	1000	0.50	7.59	83.3	0.30	0.01	3.75	0.42	95.9	1.65	10.4	79.9	0.35	<0.01	3.55	0.59	96.5
Oral (multiple dose; radiolabeled EPG on Day 15)	1000	1.68	5.38	78.3	0.52	<0.01	3.34	0.36	89.6	0.89	6.86	80.7	0.24	<0.01	1.42	0.17	90.3
Oral (single dose)	3000	0.58	5.90	80.1	0.25	<0.01	5.31	0.60	92.7	1.26	9.92	77.1	0.35	0.01	2.84	0.52	92.0

<sup>a</sup>Includes cage wash/wipe with 1% trisodium phosphate and hexane.

<sup>b</sup>Ethoxyethanol:ethanolamine trap; includes backup.

#### 4.2.3 Metabolism and Disposition of EPG-14 Oleate in Rats (HWI 6226-109; E-014)

The purpose of this study was to assess the bioavailability and extent of absorption, distribution, elimination, and biotransformation of oxypropylene-<sup>14</sup>C-EPG-14 oleate in CrI:CD<sup>®</sup>BR rats (110 to 189 g). Forty-four treated animals were divided into five groups. A preliminary group of 4 animals (2/sex) was dosed to determine whether expired air and organic volatiles needed to be collected from subsequent groups. The remaining 40 animals were divided into 4 groups of 10 animals (5/ sex): single intravenous low dose group at 30 mg/kg, single oral low dose group at 1,000 mg/kg, multiple oral low dose at 1,000 mg/kg (14 daily non-radiolabeled doses followed by a single radiolabeled dose on the 15<sup>th</sup> day), and single oral high dose group at 3,000 mg/kg. All groups had urine, feces, expired carbon dioxide (CO<sub>2</sub>), and organic volatiles collected. The animals were sacrificed 7 days after the administration of the radiolabeled dose and various tissues<sup>5</sup> and blood were collected and analyzed for total radioactivity. Mean recovery of radioactivity in this study is summarized in Table 4-3.

The majority of the radioactivity was found in the feces in all oral dose groups, with values ranging from 79.7% to 89.4% of the dose. A relatively minor amount of the dose was excreted in the urine with values ranging from 1.73% to 6.80% of the dose. The other significant route of elimination was through CO<sub>2</sub>, which represented from 3.11% to 7.15% of the dose. The excretion pattern after i.v. administration was different than that observed for the oral groups. Urine and CO<sub>2</sub> were the predominant means of excretion, with 34.4% and 37.3% of the dose excreted in urine, and 41.8% and 36.7% of the dose excreted as CO<sub>2</sub> by males and females, respectively. Feces represented 11.2% and 13.5% of the dose for males and females, respectively.

Tissue residue levels were in general very low after oral administration. The tissues that contained the highest concentrations of radioactivity in all dose groups were liver, kidney, spleen, and lungs. The majority of the dose after oral administration passed unabsorbed through the gastrointestinal tract and into the feces where it was recovered. Parent EPG-14 oleate was the major component of the feces radioactivity, although the mono- and diester were also found indicating that degradation of EPG's are occurring in the gastrointestinal tract. The highest mean percentage of radioactive dose recovered in tissues after i.v. administration was in the residual carcass for males 3.20% and in the liver for females 4.09%. For males, the liver contained the next highest mean percentage of dose with 2.71%. For females, the residual carcass contained the next highest mean percentage of dose with 2.36%.

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<sup>5</sup> Tissues included bone (femur), brain, fat, heart, kidneys, large intestine, liver, lungs, muscle (thigh), ovaries, small intestine, spleen, stomach, tail, testes and uterus.

**Table 4-3 Mean recovery (%) of radioactivity from <sup>14</sup>C-PGU-EPG-14 oleate in rats (5/group) after 7 days**

Route	Dosage (mg/kg bw)	Males								Females							
		Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total	Cage wash <sup>a</sup>	CO <sub>2</sub> <sup>b</sup>	Feces	Urine	Volatiles	Carcass	Tissues	Total
Intravenous (single dose)	30	0.31	41.8	11.2	34.4	0.03	3.20	5.11	96.1	0.93	36.7	13.5	37.3	0.04	2.36	5.59	96.5
Oral (single dose)	1000	0.28	3.11	89.4	1.73	<0.01	0.26	0.06	94.9	1.07	4.88	79.7	5.86	0.01	0.29	0.07	91.9
Oral (multiple dose; radiolabeled EPG on Day 15)	1000	0.99	6.30	85.4	3.81	<0.01	0.60	0.18	97.3	1.23	7.15	84.0	4.89	0.01	0.46	0.11	97.8
Oral (single dose)	3000	0.95	5.84	86.5	5.59	<0.01	0.69	0.12	99.7	1.79	5.42	85.8	6.80	<0.01	0.55	0.11	100

<sup>a</sup>Includes cage wash/wipe with 1% trisodium phosphate and hexane.

<sup>b</sup>Ethoxyethanol:ethanolamine trap; includes backup.

## 5.0 PRECLINICAL SAFETY

Preclinical studies were conducted with H-EPG-05 HR/SO 9:1 unless otherwise stated. EPG-05 HR/ST 45:55 is a somewhat softer version at average normal body temperature and was selectively investigated in safety studies. It is worthwhile to note that, unlike olestra, EPG is not strongly hydrophobic and exhibits far less interaction with fat-soluble substances including fat soluble vitamins. As such, vitamin fortification of animal diets was not required in any of the EPG preclinical safety studies including lifetime studies in rats and mice as well as up to 3 generations in reproductive and development studies. This differed from studies conducted with olestra, which required vitamin fortification. It is also important to note that residues of EPG were not found in any tissues from any animals placed on study, indicating efficient clearance and absence of accumulation even following lifetime administration.

### 5.1 Overview of Preclinical Safety

A battery of preclinical feeding studies was initiated to assess the safety of the core compound, H-EPG-05 HR/SO 9:1, including carcinogenic activity and the potential to cause developmental anomalies in several animal species. In addition, a series of mutagenicity studies were conducted with H-EPG-05 HR/SO 9:1, as well as other EPG versions (e.g., H-EPG-05 soyate and H-EPG-14 soyate).

The preclinical studies showed no adverse treatment related changes to the general health and appearance of the animals, or on the conventional parameters measured including but not limited to: growth, feed consumption, body weight, clinical chemistry, hematology, reproductive performance and fetal development. Feeding EPG to rats, mice, rabbits, dogs and micro-pigs produced no observed adverse findings in the GI tract structure or function. Minor fluctuations in fat soluble vitamin status were evident in preclinical studies, however, the concentrations of fat soluble vitamins in the liver (e.g., vitamins A and E) and serum (e.g., vitamin D) remained within the historical limits of species traditionally used in animal studies involving lifetime dietary exposures. Refer to Section 7.2 for additional information concerning the effect of EPG on fat-soluble vitamin status.

Articles for the following preclinical studies have been published in *Regulatory Toxicology and Pharmacology*: (i) mutagenicity assays (Section 5.8); (ii) 90-Day Dietary Safety Study with Esterified Propoxylated Glycerol (EPG) in Rats (Section 5.2.1); (iii) 90-Day Safety Study of Esterified Propoxylated Glycerol (EPG) Administered in the Feed to Yucatan Micro-Pigs® (SUS SCROFA) (Section 5.2.4); (iv) One Generation Reproduction Study of Esterified Propoxylated Glycerol (EPG) Administered in the Feed to CD® (Sprague-Dawley) Rats (Section 5.6.1); and

(v) Developmental Toxicity Evaluation of Esterified Propoxylated Glycerol [H-EPG-05 HR/SO (9:1)] Administered in the Diet to New Zealand White Rabbits (Section 5.3).

**5.1.1 Determination of the Homogeneity and Stability of Test Diets for Dietary Safety Studies with Esterified Propoxylated Glycerol (EPG) in Rodents E-057 (HWI 6226-124; E-057)**

This study was conducted in order to establish homogeneity and stability of the prepared test diets for the following studies: (i) “Combined Chronic Dietary Safety Study and Carcinogenicity Study with Esterified Propoxylated Glycerol (EPG) in Rats” (HWI 6226-120); (ii) “Combined Chronic Dietary Safety Study and Carcinogenicity Study with Esterified Propoxylated Glycerol (EPG) in Mice” (HWI6226-121); (iii) “90-Day Dietary Safety Study with Esterified Propoxylated Glycerol (EPG) in Rats” (HWI 6226-122); and (iv) “90-Day Dietary Safety Study with Esterified Propoxylated Glycerol (EPG) in Mice” (HWI 6226-123). These studies will be referred to as the dietary EPG safety studies.

Stability of EPG and  $\alpha$ -tocopherol was determined from five sets of samples of test diet (a set of samples consisted of two approximately 100 g samples). One set of samples was analyzed on the day after diet preparation; the second set was stored 5 days refrigerated ( $5^{\circ} \text{C} \pm 3^{\circ}$ ), then 9 days at room temperature and then an additional 7 days refrigerated (a total of 12 days refrigerated and 9 days at room temperature); the third set was stored for 3 weeks at room temperature; the fourth set was stored for 8 weeks refrigerated and then 11 days at room temperature; and the fifth set was stored for 7 weeks in a freezer set to maintain  $-20^{\circ} \text{C} \pm 10^{\circ}$ . When the storage periods for each set of samples was completed, one sample from each set was analyzed for  $\alpha$ -tocopherol content, and the other sample from each set was analyzed for EPG content. To evaluate stability of each test diet, analytical results of samples from the four types of storage conditions were compared with results of analyses done on the day of diet preparation.

Results of the analysis of stability samples for  $\alpha$ -tocopherol and EPG content are shown in Table 5-1.

**Table 5-1 Summary of results of stability analyses**

Storage Conditions	$\alpha$ -Tocopherol (% of initial levels)	EPG Concentration (% , mean of replicate analyses)
Refrigerated for 12 days & 9 days at room temperature	94.5 – 102	98.8 – 99.6
Room temperature for 3 weeks	89.0 – 94.3	95.8 – 96.4
Frozen for 7 weeks	103 -106	97.2 – 98.7
Refrigerated for 8 weeks & 11 days at room temperature	81.6 – 90.1	93.6 – 95.9



The data from this study demonstrated that the mixing procedures used in the dietary EPG safety studies produced homogeneous diet preparations and that the diet preparations were stable under conditions used in the dietary EPG safety studies.

## **5.2 Subchronic Toxicity Studies with Esterified Propoxylated Glycerol (EPG)**

### **5.2.1 90-Day Dietary Safety Study with Esterified Propoxylated Glycerol (EPG) in Rats (Christian and Bechtel, 2014)**

The purpose of this study was to assess the subchronic toxicity of a representative version of EPG when given to Crl:CD®BRVAF/Plus® rats (Sprague-Dawley derived; approximately 36 days old; males weighed between 147 and 193 g, and females weighed between 116 and 162 g) by dietary admixture for at least 90 days. Rats (n=700) were randomly assigned to five groups (70 animals/sex/group, subdivided into subsets A through F for each sex) and administered concentration levels of 0, 0.5, 1.0, and 2.0 g EPG/kg of bw/day (g/kg/day) through adjusted diets, or a fixed intake of 5.0% (w/w) in the diet. The latter is expected to result in a decrease in EPG intake over time; the result of feed consumption in g/day remaining relatively constant and the mean body weights increasing markedly over time so that mean feed intake in g/kg/day decreases markedly over time. All diets were prepared weekly and provided *ad libitum*.

Animals were housed individually in stainless steel, screen bottom cages, and were observed twice daily (a.m. and p.m.) for mortality, moribundity, and signs of toxicity. Body weights and food consumption were recorded weekly. Ophthalmic examinations were performed on all animals before study initiation and during Week 13 for animals given 0 g/kg and 5.0% (w/w) EPG. Hematology and clinical chemistry evaluations were done on 10 animals/sex/group before sacrifice at Weeks 5 (Subset A) and 14 (Subset C). Liver vitamin A (trans-retinol) and E (ex-tocopherol) and serum vitamin D level (25-OH vitamin D, total) determinations were done on 10 animals/sex/group during Weeks 5 (Subset A, animals necropsied) and 14 (Subset D, animals discarded without necropsy). Samples of liver, kidney, spleen, and adipose tissue were collected for EPG level determinations from 10 animals/sex killed and discarded without necropsy during Weeks 5 (Subset B) and 14 (Subset F); assays were done on the tissues from controls and animals given 5.0% (w/w) EPG. Ten animals/sex/group (Subset F) were housed in metabolism cages for the collection of fecal matter for analysis of EPG, EPG metabolites, cholesterol, fatty acids, and total bile acids. In addition, ten animals/sex/group were sacrificed and subjected to pathological evaluation at Weeks 5 (Subset A) and 14 (Subset C), and 20 additional animals/sex/group (Subset E) were sacrificed at Week 14 for pathological evaluation. At scheduled necropsies, animals were anesthetized, weighed, and exsanguinated. Macroscopic observations were recorded; selected organs were weighed; and selected tissues were preserved. Microscopic examinations were performed on all Subset A, C, and E animals in the control and 5.0% (w/w) EPG groups; on the gastrointestinal tract, lungs, liver, kidneys, and

all macroscopic lesions from animals in the remaining groups; and on all animals (regardless of subset designation) that died or were sacrificed in a moribund condition.

Results of the study showed that dietary administration of EPG at levels of 0.5, 1.0 and 2.0 g/kg, or 5% (w/w) to rats for at least 13 weeks was not associated with any adverse effects. The levels of liver vitamins A and E and serum vitamin D were generally decreased in EPG-treated animals at all concentration levels. However, there was no evidence of vitamin deficiency as assessed by growth, clinical observations, clinical pathology or anatomical pathology endpoints. Prothrombin time (PT), measured as an indicator of vitamin K status, was not significantly affected. Based on the results of this study, it was not possible to establish a no-observable-effect level (NOEL). The possible effect of EPG on vitamin levels in the absence of any clinical signs of deficiency was not considered "adverse" *per se*. As such, the adjusted concentration of 2 g/kg and the fixed intake of 5% EPG (equivalent to an average EPG intake of approximately 6 g/kg bw/day in the beginning of the study and declining to approximately 2 g/kg bw/day) were considered to represent no-observable-adverse-effect levels (NOAEL).

#### **5.2.2 90-Day Dietary Safety Study with Esterified Propoxylated Glycerol (EPG) in Mice (HWI 6226-123; E-008)**

The purpose of this study was to assess the subchronic toxicity of a representative version of EPG when given to Crl:CD-1<sup>®</sup>(ICR)BR mice (approximately 7 weeks old; males weighed between 24 and 33 g and females weighed between 18 and 26 g) by dietary admixture for at least 90 days. Mice (n=800) were randomly assigned to five groups (80 animals/sex/group, subdivided into subsets A through H for each sex) and administered concentration levels of 0, 1.0, 2.0, and 5.0 g EPG/kg of bw/day through adjusted diets, or a fixed intake of 5.0% (w/w) in the diet. All diets were prepared weekly and provided *ad libitum*.

Animals were housed individually in stainless steel, screen-bottom cages, and were observed twice daily (a.m. and p.m.) for mortality, moribundity, and signs of toxicity. At least once weekly, each animal was removed from its cage and examined. Body weights and food consumption were recorded weekly. Ophthalmic examinations were performed on all animals before study initiation and during Week 13 for animals given 0 g/kg/day and 5.0% (w/w) EPG. Hematology (10 animals/sex/group) and clinical chemistry (10 animals/sex/group) evaluations were performed before the interim sacrifice at Week 6 (Subsets A and B) and the terminal sacrifice at Week 14 (Subsets D and E). Liver vitamin A and E and serum vitamin D (25-OH vitamin D, total) level determinations were performed on samples collected from 10 animals/sex/group during Weeks 6 (Subset C) and 14 (Subset G). EPG level determinations were done on liver, spleen, and kidney tissue from 10 animals/sex in Groups 1 and 5 collected during Weeks 6 (Subset A) and 14 (Subset H). Ten animals/sex/group were sacrificed and subjected to pathological evaluation at Weeks 6 (Subset B) and 14 (Subset D), and 20 additional animals/sex/group (Subsets E and F) were also sacrificed at Week 14 for pathological evaluation. At scheduled

necropsies, animals were anesthetized, weighed, and exsanguinated. Macroscopic observations were recorded, selected organs were weighed, and selected tissues were preserved. Microscopic examinations of all preserved tissue were performed on all animals in the control group and 5.0% (w/w) EPG group (Subsets B, D, E, and F). Microscopic examination of lesions, gastrointestinal tract, lungs, liver, and kidneys was also performed on all animals in the remaining groups.

Results of the study showed that dietary administration of EPG at levels of 1.0, 2.0, and 5.0 g/kg, and at 5.0% (w/w) for at least 90 days was not associated with adverse effects. Body weight gains were slightly increased over the treatment period in animals given 5.0% (w/w) EPG. The mean liver vitamin A levels at Week 14 were statistically lower for males given 5.0 g/kg and 5% EPG. For females at Week 14, liver vitamin A levels were lower for all EPG-treated groups, however, not in a dose-related manner and statistical significance was noted for only for females given 1.0 g/kg. At Week 14, concentration-related decreases in liver vitamin E were noted for both males and females exposed to the test material. At Week 14, serum vitamin D (25-OH vitamin D, total) levels in males given 5.0 g/kg and 5.0% EPG and females given 2.0 and 5.0 g/kg and 5.0% EPG were reduced; these test material-related differences were statistically significant. Males given EPG at levels of 1.0 and 2.0 g/kg and females given 1.0 g/kg had serum vitamin D levels similar to those of controls. There was no evidence of vitamin deficiency in any of the animals as assessed by growth, clinical observations, clinical pathology, or anatomical pathology endpoints. Based on these results, the NOAEL was 5.0% (w/w) EPG in the diet, equivalent to 6.2 g/kg (males) and 8.1 g/kg (females) at study termination.

### 5.2.3 Thirteen Week Safety Study of EPG Administered in the Feed to Beagle Dogs (T.P.S. Study No. 460D-502-634-92; E-009)

This study was designed to evaluate the toxicity of EPG when administered as a dietary admix to beagle dogs, daily, for at least 13 weeks. Male and female (5/sex/group) purebred beagle dogs (approximately 7 to 8 months old) were fed modified Teklad Basal Dog Diet Certified Meal supplemented with 6% corn oil which contained EPG; the experimental design of this study was as follows:

Group No.	EPG Theoretical Concentration Level*		EPG Mean Actual Dose (g/kg bw/day)*		Number of Dogs per Group	
	(g/kg bw/day)	% Diet (wt/wt)	Male	Female	Male	Female
AVK1 (Control)	0.0	0	0	0	5	5
AVK2	1.5	5	1.6	1.6	5	5
AVK3	3.0	10	3.2	3.2	5	5
AVK4	5.0	17	5.2	5.3	5	5

\* Approximate levels; animals received feed containing 5, 10 and 17% EPG, mixed accordingly throughout the study to deliver 1.5, 3.0 and 5.0 g/kg bw/day, respectively.

Dogs ate the isocaloric diets offered to them over the 2-hour daily feeding period and food consumption was measured daily. They were housed individually in adjacent runs with chain linkwire sides, epoxy coated floors, and elevated resting boards, and were observed twice daily for pharmacological, toxicological, and behavioral effects; feces were inspected for phase separation (*i.e.*, appearance of oily layers or deposits in the stools). Detailed physical examinations including ophthalmoscopy were conducted pretest, during Week 7, and just prior to necropsy. A general physical examination was done each week on each animal. Body weights were recorded pretest and weekly thereafter. Blood samples from fasted animals were obtained at pretest and during Weeks 6 and 13 for evaluation of hematologic and clinical chemistry parameters. Serum samples obtained just prior to necropsy were analyzed for total Vitamin D and 25-OH-Vitamin D concentrations. Urinalysis (including urine chemistries) and water consumption evaluations were conducted pretest and during Weeks 6 and 13/14. Fecal samples obtained during Weeks 6 and 13/14 were analyzed for total fat, total cholesterol, total fatty acids and profile, total fecal bile acid, and calcium. A separate fecal sample was obtained during Weeks 6 and 13/14 for possible gut microflora assay. At necropsy samples of liver (all lobes), kidney, spleen and adipose tissue were taken for EPG analysis; an additional liver sample (all lobes) was analyzed for vitamin A and E content. All dogs were subjected to a complete postmortem examination.

No treatment-related mortality/morbidity occurred. No consistent or distinct EPG treatment related adverse pharmacological/toxicological or behavioral effects were noted during this evaluation. Physical and ophthalmic examinations at termination indicated no treatment-related effect. The dogs ate the isocaloric diets offered to them (over each 2-hour daily feeding period) indicating EPG was palatable up to 5.0 g/kg/day in the diet. Visual inspection of the feces daily indicated no phase separation (*i.e.*, appearance of oily layers or deposits in the stools). Analysis of body weight gain, feed efficiency, water consumption, bowel transit times, hematology and serum chemistry parameters, urinalysis data (including urine chemistries), feces (for total fat, total fatty acid and calcium), organ weights (absolute, relative to body or brain weight) and vitamin A concentration of the liver did not indicate a treatment-related effect.

Results of chemical analyses of liver, kidney, spleen and adipose tissue showed that there was no EPG found in any of the tissues examined, within the limits of detection. Fecal chemical assay results for EPG and the mono and diester metabolite concentrations showed that all fecal samples from dogs receiving 17% EPG triester diet contained EPG triester, diester, and monoester after 6 weeks of dosing at levels of 11.7 to 29.2 g, 1.21 to 3.07 g, and 0.183 to 0.517 g, respectively, for males; 15.2 to 35.3 g, 1.44 to 3.61 g and 0.348 to 0.784 g of triester, diester and monoester were present in the feces from female dogs in a 24-hour period during week 6 of dosing. The amounts of EPG triester, diester, and monoester found in the feces of male dogs in a 24-hour period during week 13 of dosing were 5.08 to 37.8g, 0.494 to 3.25g and 0.121 to 0.553 g, respectively; 13.4 to 27.3 g, 1.50 to 3.03 g, and 0.321 to 0.596 g of triester, diester, and

monoester were present in the feces of female dogs from a 24-hour period during Week 14 of dosing.

Chemical analysis of feces collected during Week 6 and 13/14 indicated a treatment-related significant increase in cholesterol and decrease in total bile acid at 5 g EPG/kg/day. A significant decrease in the liver vitamin E content of the Group AVK4 animals was considered to be treatment-related. A concentration-dependent relationship was apparent with respect to the total vitamin D serum levels which were significantly reduced in all groups. The serum level of 25-OH-Vitamin D was significantly reduced at 3 and 5 g EPG/kg/day in a concentration-dependent manner. Effects of EPG on these serum vitamin levels were considered to be treatment-related. PT and activated partial thromboplastin time (APTT), measured as indicators of vitamin K status, were not significantly affected.

Gross necropsy and histological examination of all tissues indicated no distinct or consistent treatment-related effects. In summary, an increase in fecal cholesterol, a decrease in total fecal bile acid, and a decrease in vitamin E content of the liver were considered treated-related at a level of 17% EPG in the diet, or 5 g EPG/kg/day. Associated with dietary administration of EPG, the NOEL for serum vitamin D was <1.5 g EPG/kg/day, as serum vitamin D levels were significantly reduced in all groups even though there were no indications of vitamin deficiency expressed clinically. The NOEL for the biologically active 25-OH-Vitamin D metabolite was 1.5 g EPG/kg/day.

#### 5.2.4 90-Day dietary toxicity study with esterified propoxylated glycerol (EPG) in Micropigs (Wedig and Bechtel, 2014)

The subchronic (90-day) toxicity of EPG was assessed in Yucatan micropigs (approximately 8 to 10 months old). Animals (5/sex/group) received feed (Certified Agway® Prolab® Minipig Diet Meal) containing 5, 10, and 17% EPG, mixed accordingly throughout the study to deliver 1.5, 3, and 5 g/kg/day of EPG, respectively. Corn oil served as the vehicle control (0 g/kg bw/day). The study design is also summarized in Table 5-2.

**Table 5-2 EPG concentration and group composition**

Group	Treatment		Number of animals/sex
	EPG*	Dietary Concentration <sup>†</sup>	
Control (AVI1)**	0 g/kg/day	0% (w/w)	5
Low EPG (AVI2)	1.5 g/kg/day	5% (w/w)	5
Mid EPG (AVI3)	3 g/kg/day	10% (w/w)	5
High EPG (AVI4)	5 g/kg/day	17% (w/w)	5

\* Approximate levels; animals received feed containing 5, 10 and 17% EPG, mixed accordingly throughout the study to deliver 1.5, 3.0 and 5.0 g/kg bw/day, respectively.

<sup>†</sup> % (w/w) = weight of EPG per weight of basal diet including corn oil

\*\* The basic diet was supplemented with 4% (w/w) corn oil; test diets contained EPG and 4% (w/w) corn oil as vehicle

Micropigs were observed twice daily for toxicological, pharmacological, and behavioral effects. Feed consumption and dietary levels of EPG were determined on a weekly basis. Physical and ophthalmic examinations, body weights, urinalysis, hematology, clinical chemistry, water intake, bowel transit times, organ weight, organ tissue analysis for EPG, fecal assays, vitamin assays, gross necropsy, and histopathology were used to evaluate the effects of EPG.

EPG was palatable up to 5 g/kg/day in the diet. No treatment-related morbidity/mortality occurred. No consistent or distinct EPG treatment-related adverse pharmacological/toxicological or behavioral effects were noted. No treatment-related effects were observed during the physical and ophthalmic examinations. Analysis of body weight gain, feed efficiency, water consumption, bowel transit times, hematology and serum chemistry parameters, urinalysis data, feces, and organ weights indicated no treatment-related effects. Chemical analysis of liver, kidney, spleen, and adipose tissue yielded negative data for EPG residue. Gross necropsy and histopathology examinations indicated no treatment-related effects.

PT and APTT, measured as indicators of vitamin K status, were not significantly affected. EPG significantly affected liver vitamin A and serum vitamin D. A significant decrease in the liver vitamin A content was observed in animals fed 5 g of EPG/kg/day. EPG demonstrated a concentration-dependent effect on the levels of total vitamin D and the biologically active vitamin D metabolite, 25-OH-vitamin D. Specifically, total vitamin D serum levels were significantly reduced in all groups, while serum levels of 25-OH-vitamin D were significantly reduced in animals administered 3 or 5 g of EPG/kg/day. Although a NOEL for effects of dietary EPG on total vitamin D serum levels was not established, a NOEL for effects on 25-OH-vitamin D levels was determined to be 1.5 g of EPG/kg/day, or 5% dietary EPG concentration.

### **5.3 Subchronic Toxicity Study with EPG-05 HR/ST 45:55**

Note that the EPG version tested in this study is somewhat softer than the “core” version tested in other studies.

#### **5.3.1 90-Day Dietary Toxicity Study of EPG-05 HR/ST (45:55) in Rats (MPI Study Identification No. 728-006; E-045)**

In a 90-day toxicity study, CrI:CD<sup>®</sup>BR (VAF/Plus) rats (20/sex/group; approximately 4 weeks of age) were administered EPG-05 HR/ST (45:55) at concentration levels of 0, 0.5, 1.0, or 2.0 g/kg bw/day through adjusted diets, or a fixed intake of 5.0% (w/w) in the diet. Rats were individually housed in suspended stainless steel cages with wire-mesh floors and were observed twice daily for mortality, morbidity, and signs of toxicity. Detailed observations, body weight, food consumption, and EPG-05 HR/ST (45:55) consumption were monitored weekly. Individual body weight and an ophthalmoscopic examination were conducted before dosing. Ophthalmoscopic examination, clinical pathology assessment, and serum sample evaluation were conducted in

the last week of the study. Necropsies were performed at study termination. Results showed that serum total vitamin D (25-OH D<sub>2</sub> plus 25-OH D<sub>3</sub>) levels decreased in a dose-related manner for all groups of males and females administered EPG-05 HR/ST (45:55), and were statistically significant in the 2.0 g/kg/day and 5.0% (w/w) groups. The mean liver vitamin E concentration was statistically significantly decreased in all groups of males and females receiving EPG-05 HR/ST (45:55) in the diet when compared to controls. In females, the decreases were dose-related. The mean concentration of liver vitamin A in males was statistically significantly decreased in the 1.0 and 2.0 g/kg/day and the 5% (w/w) groups, although the decreases were not dose-related. No effect on liver vitamin A levels were noted in males receiving 0.5 EPG-05 HR/ST (45:55) g/kg/day. In females, the mean liver vitamin A concentrations were decreased in all groups administered EPG-05 HR/ST (45:55) in the diet. The decreases were not dose-related, but were statistically significant in the 0.5 and 2.0 g/kg/day and the 5.0% (w/w) females. No adverse clinical signs, gross, or microscopic pathologic evidence of vitamin A, E, or D deficiency were observed. PT and APTT, measured as indicators of vitamin K status, were not significantly affected.

#### **5.4 Chronic Toxicity Studies with Esterified Propoxylated Glycerol (EPG)**

##### **5.4.1 Combined Chronic Dietary Safety Study and Carcinogenicity Study with Esterified Propoxylated Glycerol (EPG) in Rats (HWI 6226-120; E-001)**

The purpose of this study was to assess the potential chronic toxicity and carcinogenicity of the test material, EPG, when fed to CrI:CD@BR VAF/Plus® rats (Sprague-Dawley derived) for at least 104 weeks. Rats (n=1,400) were randomly assigned to five groups (140 animals/sex/group) and administered EPG at concentration levels of 0, 0.5, 1.0, or 2.0 g/kg bw/day through adjusted diets, or a fixed intake of 5.0% (w/w) in the diet. All diets were prepared weekly and available *ad libitum*. Fifty rats/sex/group were designated for interim sacrifice and were terminated after 52 weeks of treatment.

Animals were housed individually in stainless steel, screen-bottom cages and were observed twice daily (a.m. and p.m.) for mortality, moribundity, and signs of toxicity. Body weights were recorded on the first day of treatment, weekly for 16 weeks and every four weeks thereafter; food consumption was recorded weekly for 16 weeks and every 4 weeks thereafter. Ophthalmic examinations were performed on all animals before study initiation and during Weeks 13, 26, 39, 54, 65, 78, and 91 for animals in the control and 5.0% EPG groups. Ophthalmic examinations were also done for all animals scheduled for interim (Week 52) and terminal (Week 104) sacrifice. Hematology evaluations were done on 20/sex/group at Weeks 26 and 78, and hematology and clinical chemistry evaluations were done for 20/sex/group before the interim sacrifice (Week 53/54) and terminal sacrifices (Week 105/106). Liver vitamin A, vitamin E, and serum vitamin D (25-OH vitamin D, total) level determinations were done for

10/sex/group necropsied at the interim and terminal sacrifice. Ten animals/sex/group were housed in metabolism cages for the collection of fecal matter for analysis of EPG, EPG metabolites, cholesterol, fatty acids, and total bile acids. Forty rats/sex/group were sacrificed and subjected to pathological evaluation at the Week 53/54 interim sacrifice; and all surviving animals were sacrificed at the Week 105/106 terminal sacrifice. Microscopic examinations were performed on the animals in the control and 5.0% EPG groups; on all animals that died or were sacrificed; and on the gastrointestinal tract, lungs, liver, kidney, and all macroscopic lesions from animals in the remaining groups.

Results of the study showed that dietary administration of EPG to rats at concentration levels of 0.5, 1, and 2 g/kg, or 5.0% (w/w) in the diet for at least 104 weeks was not associated with adverse effects nor were there neoplasms whose incidence suggested an association with EPG administration. Body weights were slightly increased in males given 1 g/kg, 2 g/kg, and 5.0% EPG, and in females given 5.0% EPG. The levels of liver vitamins A and E and serum vitamin D (25-OH vitamin D, total) were generally decreased in EPG-treated animals. However, there was no evidence of vitamin deficiency as assessed by growth, clinical observations, clinical pathology or anatomical pathology endpoints. PT, measured as an indicator of vitamin K status, was not significantly affected. The increased incidence of palpable masses in females given 1 g/kg, 2 g/kg and 5.0% EPG which correlated with higher incidences of mammary fibroadenomas and increased incidence of thyroid C-cell adenomas, were not considered important because these are commonly occurring spontaneous neoplasms in female rats and furthermore, the incidence at which they were observed was comparable to that observed in control animals of previous studies. Based on the results of this study, the NOAEL was 5.0% (w/w) in the diet, equivalent to 2 g/kg/day at study termination.

#### **5.4.2 Combined Chronic Dietary Safety Study and Carcinogenicity Study with Esterified Propoxylated Glycerol (EPG) in Mice (HWI 6226-121; E-002)**

The purpose of this study was to assess the chronic toxicity and potential carcinogenicity of the test material, EPG, when fed to CrI:CD-1® (ICR) BR VAF/Plus® mice (51 (males) or 52 (females) days old; males weighed between 20.4 and 33.7 g, and females weighed between 18.9 and 27.7 g) for at least 104 weeks. Mice (n=1,500) were randomly assigned to five groups (150 animals/sex/group) and administered EPG at concentration levels of 0, 0.5, 1.0, or 2.0 g/kg bw/day through adjusted diets, or a fixed intake of 5.0% (w/w) in the diet. All diets were prepared weekly and available *ad libitum*. Fifty mice/sex/group were designated for interim sacrifice and were terminated after 52 weeks of treatment.

Animals were housed individually in stainless steel, screen-bottom cages, and were observed twice daily (a.m. and p.m.) for mortality, moribundity, and signs of toxicity. Body weights and food consumption were recorded weekly for 16 weeks and at least every 4 weeks thereafter. Body weights were also recorded at the time of death for all animals that died or were sacrificed



at an unscheduled interval. Ophthalmic examinations were performed on all animals before study initiation, on all animals scheduled for interim and terminal sacrifice (Weeks 52 and 104) and on all animals in Groups 1 and 5 (control and 5.0% EPG) during Weeks 13, 26, 39, 55, 77, and 91. Hematology evaluations were done on 10 animals/sex/group at Weeks 27 and 78 and after 104 weeks; hematology and clinical chemistry evaluations were done on 10 animals/sex/group before the interim sacrifice (Week 53). Liver vitamin A, vitamin E, and serum vitamin D (25-OH vitamin D, total) level determinations were done for 10 mice/sex/group necropsied at the interim and terminal sacrifices. Ten animals/sex/group were housed in metabolism cages for the collection of feces for analysis of EPG, EPG metabolites, cholesterol, total fatty acids, and bile acids; collections were initiated during Weeks 1, 4, 13, 26, 52, 78, and 103. During these collections, body weight, food consumption, and fecal output measurements were performed daily. At scheduled necropsies, animals were fasted overnight, anesthetized, weighed, exsanguinated, and necropsied. Macroscopic observations were recorded; selected organs were weighed and selected tissues were preserved. Microscopic examinations were performed on the animals in the control and 5.0% EPG groups; on all animals that died or were sacrificed in a moribund condition; and on the gastrointestinal tract, lungs, liver, kidneys, and all macroscopic lesions from animals in the remaining groups.

Based on the results of the study, dietary administration of EPG at concentration levels of 1.0, 2.0, and 5.0 g/kg, or 5.0% (w/w), for at least 104 weeks was associated with no adverse effects. In addition, since there were no statistically significant differences in the incidence of neoplasms between the control and treated groups, EPG and was not carcinogenic. Body weights were increased slightly over the treatment period which may be related to slight increases in food consumption. At Week 53, mean liver vitamin A levels were lower in males and females given 2.0 and 5.0 g/kg, and 5% EPG compared with those of the control group; these differences were not statistically significant but were dose-related for males. Mean liver vitamin A levels at week 106 were lower for animals given all dose levels of EPG when compared to control animals. For animals given 5.0% EPG, liver vitamin A levels were reduced by approximately 16% for males and 24% for females; only the value for females was statistically significant. Group mean liver vitamin E levels were similar for control and EPG-treated males at all dose levels at Week 53. For EPG-treated females, liver vitamin E levels were reduced in a dose-dependent manner; these differences were statistically significant for animals given 5.0 g/kg and 5.0% EPG. At Week 106, mean liver vitamin E levels were reduced in a dose-dependent manner for males and females. The differences were statistically significant for males given 5.0 g/kg and for females given 2.0 or 5.0 g/kg or 5.0% EPG. There was no effect of EPG administration on serum levels of vitamin D in males or females. Despite reductions in liver vitamin A and E levels compared to control values, there was no evidence of vitamin deficiency as assessed by growth, clinical observations, clinical pathology, or anatomical pathology endpoints. The increased incidence and severity of skin lesions in males given 5.0% (w/w) EPG were associated with the clinical observation of an increased incidence of oily hair coat; this finding is

likely secondary to dermal contact with the diet (oily consistency) and is not considered a adverse effect of the test material. Based on these results, the NOAEL was 5.0% (w/w) in the diet, equivalent to 2 g/kg/day at study termination.

## **5.5 Chronic Toxicity Studies with H-EPG-05 HR/SO (9:1)**

### **5.5.1 A One-Year Chronic Safety Study of H-EPG-05 HR/SO (9:1) Esterified Propoxylated Glycerol (EPG) Administered in the Feed to the Beagle Dogs (MPI Study Identification No. 728-001)**

Forty purebred beagle dogs (approximately 5-6 months of age) were fed EPG at levels of 0, 1, 2, and 3 g/kg bw/day in Modified Teklad Basal Mix for Dog Diet #7058 supplemented with 6% (w/w) corn oil (5/sex/group), for one year in order to evaluate potential effects attributed to long-term EPG consumption. Animals were individually housed in standard nonspecialized stainless steel dog cages and were observed for mortality, signs of gross toxicity, clinical signs, body weights, food consumption/food efficiency, and water consumption, as well as electrocardiogram, ophthalmoscopic, hematologic and urinalysis findings. EPG concentrations were measured in adipose tissue, liver, kidney, and spleen samples. Additional tests were conducted to measure EPG its metabolites, as well as fatty acids in the feces, and recovery of EPG and its breakdown products in feces. Other parameters evaluated included: bowel transit times, vitamin concentrations in plasma, serum, and liver tissue, and mineral concentrations in liver tissue and bone. Necropsies were performed on all animals.

Effects attributed to EPG consumption included statistically significant increases in fecal fat and fatty acids in the 3 g/kg bw/day group, statistically significant decreases in fecal calcium in the 3 g/kg bw/day group, and statistically significant decreases in serum phosphorus in the 3 g/kg bw/day females at 3 and 6 months, and in the 1 g/kg bw/day females at 6 months. All treated groups, males and females, had decreases in serum vitamin E concentrations that were statistically significant at 3 g/kg bw/day in males, and 2 and 3 g/kg bw/day in females. The average concentrations of iron and vitamin E in liver tissue were found to be statistically significantly lower for males in all groups compared to controls. Additionally, bone zinc was statistically significantly decreased at 2 and 3 g/kg bw/day in males, but was increased compared to controls at 1 g/kg body weight/day. PT and APTT, measured as indicators of vitamin K status, were not significantly affected. Intact EPG was not found at the level of detection in any tissues analyzed and no clinical signs of vitamin deficiency were observed in this study.

### **5.5.2 A One-Year Chronic Safety Study of Esterified Propoxylated Glycerol H-EPG-05 HR/SO (9:1) (EPG) Administered in the Feed to Yucatan Micropigs® (Sus scrofa) (MPI Study Identification 728-002; EPG-041)**

The objective of this study was to evaluate the effects of H-EPG-05 HR/SO (9:1) administration in feed to Yucatan Micropigs® (Sus scrofa) for one year. Yucatan Micropigs® (approximately 9 months of age) were fed Certified Agway® Prolab® Minipig Diet Meal supplemented with 4% corn oil containing H-EPG-05 HR/SO 9:1 daily. Animals received 0, 1, 2, or 3 g/kg/day of H-EPG-05 HR/SO 9:1 through the diet (5/sex/group). All groups were fed H-EPG-05 HR/SO 9:1-containing micropig feed in the morning and at night. In addition, animals were fed 100 g/day of feed (containing no EPG) at noon to provide additional calories; this amount was increased to 250 g/day starting Week 47. Animals were individually housed in pens consisting of solid floors and wire sides or chain link fencing as dividers, with laboratory grade ASPEN wood shavings. Feed intake, EPG consumption, water intake, physical and ophthalmic examinations, clinical and behavioral observations, body weight gain, hematology, serum chemistry, urinalysis, bowel transit times, organ weights, necropsy, histopathology, electrocardiograms, fecal assays, tissue assays for H-EPG-05 HR/SO 9:1, vitamin and mineral assays, and bone density, bone mineral content, bone mineral density, and tissue mass measurements were used to evaluate the effects of H-EPG-05 HR/SO 9:1.

Effects related to H-EPG-05 HR/SO 9:1 included decreased mean body weights (referable to caloric restriction, not toxicity) in all groups of treated males, increased fecal fatty acids in pigs administered 3 g/kg/day (referable to H-EPG-05 HR/SO 9:1 and its breakdown products), and decreased mean total serum 25-hydroxy vitamin D concentrations in all treated groups. Although a decrease in mean total serum 25-hydroxy vitamin D concentrations was noted in all treated groups, there were no clinical signs of a vitamin deficiency as evidenced by the lack of changes in bone density, bone mineral content, and bone mineral density upon measurement with a X-ray densitometer. No other parameters including PT and APTT, which were measured as indicators of vitamin K status, were affected by H-EPG-05 HR/SO 9:1 consumption. High performance liquid chromatography (HPLC) analysis confirmed that intact H-EPG-05 HR/SO 9:1 was not present in liver, kidney, spleen, or body fat.

## **5.6 Reproductive and Developmental Toxicity**

### **5.6.1 One-Generation Reproduction Study of Esterified Propoxylated Glycerol (EPG) Administered in the Feed to CD® (Sprague-Dawley) Rats (Tyl and Bechtel, 2014a)**

This study investigated the reproductive effects following continuous exposure of Crl:CD® (SD)Br rats (approximately 6 weeks old; mean male weight  $183.9 \pm 1.1$  g, mean female weight  $151.5 \pm 1.0$  g) to EPG in the diet (30 animals/sex/group) at 0.0% (group 0), target levels of 0.5% g/kg/day (group 1), 1.0 g/kg/day (group 2), and 2.0 g/kg/day (group 3), and fixed 5.0% EPG

(w/w) (group 4), all in 6% corn oil (vehicle). Dietary concentrations of EPG for groups receiving 0.5, 1.0, and 2.0 g/kg/day of test material were adjusted weekly to maintain target EPG intake, throughout the prebreed period. Animals were exposed for a 13-week prebreed period, and through two breeding cycles for F0 parental animals, and up to postnatal day 91 for F1a and F1b offspring. Results from this study aided in the design of a subsequent three-generation reproductive toxicity study (refer to Section 5.6.2).

Parameters examined included, body weights, weight gains, feed consumption, clinical signs, reproductive indices and offspring litter sizes, pup survival and body weights, and histopathology of parental reproductive organs. The study also examined possible effects on blood clotting, parental and offspring immunologic status, histopathology of organs related to immunological function, neurological effects in parents and offspring, developmental effects in offspring, liver and serum fat-soluble vitamin status, and the possible presence of the test material in selected organs of parental and offspring animals.

Results indicated that dietary administration of EPG at levels of 0.5, 1.0, and 2.0 g/kg, and 5% (w/w) to rats for at least 13 weeks was not associated with adverse effects, except for that on liver vitamin status. Vitamin E levels exhibited concentration-related statistically significant reductions in all evaluated groups, with the exception of F1b(A) male weanlings and satellite group F1b(B) males and females. There was no evidence of vitamin D deficiency except in F0 parental, F1a(A) and F1b(A) weanling females, and no evidence of vitamin A deficiency except in F0 parental, F1a(C) and F1b(C) postnatal day 91 females. There were no effects on reproduction of the F0 parental animals for either F1a or F1b mating; evidence for dystocia was present in all groups, including the vehicle control group, with no concentration-dependent response pattern. No treatment-related effects on postnatal growth or development (physical or behavioral), immunological status, blood clotting, and parental general status were observed. EPG was not detected in any of the 360 liver samples from the high concentration and control groups. With respect to kidney and spleen samples, there were two and seven positive samples, respectively, out of 360 total samples for each organ. According to the authors, the positive samples were not the result of contamination during necropsy or analyses, were evenly divided between high concentration and control animals, and were not associated with other measures indicative of *in vivo* systemic exposure.

Based on the results of this one-generation reproduction study, the NOAEL was 5.0% EPG. Also, in the absence of any effects on behavioral development, immunologic status, and blood clotting, and with group 4 animals tolerating a fixed dietary EPG percentage, it was recommended that the three-generation study with two litters per generation utilize fixed dietary percentages with the highest concentration 5.0% EPG, and endpoints examined not include behavioral, immunologic, or coagulation assessments.

**5.6.2 Three Generation Reproduction Study (with a Teratology Phase) of Esterified Propoxylated Glycerol (H-EPG-05 HR/SO (9:1); EPG) Administered in the Feed to CD<sup>®</sup>(Sprague-Dawley) Rats (RTI Study Identification No. 65C-5304-02; E-003)**

To evaluate the potential effects of EPG on reproduction and development, in three generations of 240 CrI:CD<sup>®</sup> (SD)Br rats (approximately 6 weeks old; mean male weights 182.0-184.6 g and mean female weights 151.8-154.6 g) were fed EPG in a modified NIH-07 diet *ad libitum*, 7 days/week, at concentration levels of 0, 1, 2, and 5%, in 6% corn oil vehicle (30 animals/sex/group). The parental generation (designated F0, F1a and F2a) was exposed over a 10-week pre-breed, 3-week mating, 3-week gestation, and 3-week lactation period. The study design is summarized in Table 5-3.

**Table 5-3 Summary of study design for three-generation reproduction study with H-EPG-05 HR/SO (9:1)**

Groups	No. of Animals	Animal Fate	Endpoints Assessed
F0, F1a parental animals <sup>6</sup>	30/sex/group	Males sacrificed at end of second mating  Non-pregnant females sacrificed at least 3 weeks after last day of second cohabitation period  Parent females sacrificed at weaning of each F1b litter or F2b litter	Mortality, clinical observations, reproductive and lactational indices, body weights, feed consumption, gross lesions, histopathology exams on animals in the high concentration and control groups only
F2a parental animals	30/sex/group	Males sacrificed at end of second mating  Non-pregnant females sacrificed at least 3 weeks after last day of second cohabitation period  Parent females were sacrificed on gestational day 20	Mortality, clinical observations, reproductive and lactational indices, body weights, feed consumption, gross lesions, histopathology exams on animals in the high concentration and control groups only; Reproductive organs from males and females that failed to mate; uterus plus one attached ovary in dams to examine ovarian corpora lutea and uterine contents
F1a, b offspring F2a, b offspring F3a, b offspring	F1a,b and F2a,b culled to 10 pups per litter on pnd 4  F1a and F2a pnd 21 weanlings assigned as parents for next generation  F3a and F3b 10/sex/group	F1b, F2b, F3a sacrificed at weaning  F3b fetuses part of teratology phase; c-sectioned from F2a dams on gestational day 20	Body weights, clinical observations, survival on pnd 4, 7, 14 & 21, body weights recorded on pnd 0  All live pups were counted, sexed, and examined grossly on pnd 0, 4, 7, 14 & 21  F3b animals were measured for crown rump length, morphological abnormalities including cleft palate; external, skeletal, visceral exams performed
F1a(A) satellite	10/sex/group	Animals sacrificed at weaning	Body weights, clinical observations and liver weights
F1b(A) satellite	10/sex/group	Animals were sacrificed at weaning	Body weights, clinical observations and liver weights
F2a(A) satellite	10/sex/group	Animals were sacrificed at weaning	Body weights, clinical observations and liver weights
F2b(A) satellite	10/sex/group	Animals were sacrificed at weaning	Body weights, clinical observations and liver weights
F3a(A) satellite	10/sex/group	Animals were sacrificed at weaning	Body weights, clinical observations, total and differential leukocyte counts, total protein, albumin, and serum, albumin/globulin ratio, and liver weights
F3a(B) satellite	10/sex/group	Animals were sacrificed at weaning	Body weights, clinical observations and liver weights

<sup>6</sup> The parental generation designated as F0, F1a, F2a were given 0, 1.0, 2.0, and 5.0% EPG in 6% corn oil vehicle in the diet during a 10 week pre-breed period.

Animals were individually housed upon the initiation of the treatment period in solid bottom polycarbonate cages with stainless steel wire lids. Study animals were housed two per cage (one male:one female from the same dose level) during the mating periods. Females were caged separately and individually once they were successfully mated (or at the end of the mating periods). Females were housed individually with their litters during the lactation periods.

Endpoints assessed for the parental generation included mortality and clinical exams, body weights, feed consumption, complete necropsy with special attention to reproductive organs, and histopathological exams on animals in the high concentration and control groups.

F1a,b, F2a,b, and F3a,b were examined as soon as possible after birth to determine the number of viable and stillborn members of each litter. Where appropriate, litters were evaluated for survival on postnatal day (pnd) 4, 7, and 14, and at weaning (pnd 21). Litter weights were measured at the time of parturition, and pnd 4, 7, 14, and 21. On pnd 4, the size of each litter was adjusted by eliminating extra pups by random selection to yield 10 pups per litter.

F3b fetuses were c-sectioned from F2a dams on gestational day 20 and measured for crown rump length, morphological abnormalities including cleft palate; external, skeletal, and visceral malformations and variations.

Liver weights, body weights, feed consumption, and clinical observations were collected on satellite group animals F1a,b (A), F2a,b (A), and F3a,b (A). In addition, F3a (A) animals had total and differential leukocyte counts, total protein, albumin, and serum albumin/globulin ratios evaluated.

There were no treatment-related deaths for the parental animals. Clinical observations indicated no treatment-related findings throughout the study. All reproductive and lactational indices were equivalent for all matings throughout the study, and at necropsies, there were no treatment-related gross lesions.

In conclusion, continuous dietary exposure to EPG at all concentration levels through three generations resulted in no indications of systemic, reproductive, developmental or postnatal toxicity.

### **5.6.3 Developmental Toxicity Evaluation of Esterified Propoxylated Glycerol (EPG) Administered in the Diet to New Zealand White Rabbits (Tyl and Bechtel, 2014b)**

Seventy-two female New Zealand White rabbits (18/group) were fed EPG [0.0, 2.5, 5.0, and 10.0% (w/w)] in Modified Purina Certified Rabbit Chow #5322, supplemented with 6% corn oil (w/w), for 26 days (day -7 through gestational day 19) to assess effects of EPG on the developing conceptus.

All maternal animals were observed for mortality, signs of gross toxicity, clinical signs, body weights, food consumption, and gestational parameters. A significant concentration-related downward trend was observed for "maternal" weight change only for day -7 to d 0 (the first week of dietary exposure, prior to insemination) with no significant pairwise comparisons to the concurrent control group. For "maternal" feed consumption (in g/kg/day), significant concentration-related downward trends were observed for day -14 to -13, day -11 to -10 and day -14 to -7, with no significant pairwise comparisons, and all intervals prior to the initiation of administration of EPG (which began on day -7). At necropsy, all fetuses were dissected from the uterus and examined for skeletal malformations or variations, body weights, and crown-rump length. No evidence of maternal or developmental toxicity was found in this study. A NOAEL of 10% EPG (approximately 4.76 g/kg body weight/day), the highest dose tested for both maternal and developmental toxicity is proposed based on the results of this study.

## **5.7 Irritation and Sensitization Studies**

### **5.7.1 Primary Skin Irritation Study in Rabbits, Primary Eye Irritation Study in Rabbits (Hill Top Biolabs Report No. 87-1064-21 (A); E-060)**

This study was conducted to evaluate the potential of EPG-08-oleate (lot # 606617 THPG08), to produce irritation to the skin and eye of New Zealand White rabbits. The study was conducted in compliance with the conditions specified in the Regulation for the Enforcement of the Federal Hazardous Substances Act (16 CFR 1500).

For the primary skin irritation study in rabbits, the test material, a clear light yellow liquid, was applied undiluted to a one-inch by one-inch square surgical gauze patch. The patch was then applied to an intact skin area and an abraded skin area on six young adult rabbits (3M and 3F). Each animal received 0.5 mL of test material at each application site. Rabbits were housed individually in wire mesh suspension cages and allowed Purina Laboratory Rabbit Chow and tap water *ad libitum*. They were maintained on a 12-hour light/12-hour dark cycle. At the end of the 24-hour exposure period, the patches were removed and the sites were scored for erythema and edema and checked for tissue damage according to the method of Draize (1959). Two days later (72-hour reading), sites were again scored for erythema and edema and checked for tissue damage. Results from the Primary Irritation Index were found to be 1.0 based on erythema and edema. No evidence of tissue damage was found. In conclusion, the test material is not classified as a primary irritant or as corrosive following dermal application.

For the primary eye irritation study in rabbits, the test material, a clear light yellow liquid, was applied undiluted to the right eye of each of six New Zealand White rabbits (3M and 3F). It should be noted that these are not the same rabbits as used in the aforementioned primary skin irritation study. Each animal received 0.1 mL of the test material. The eye was not rinsed following the application. Rabbits were housed individually in wire mesh suspension cages and



allowed Purina Laboratory Rabbit Chow and tap water *ad libitum*. They were maintained on a 12-hour light/12-hour dark cycle. The eyes of each rabbit were examined approximately 24 hours prior to treatment to assure that they had no pre-existing lesions which could compromise the study. The eyes were graded for corneal changes, conjunctival changes, and changes in the iris approximately 24 hours following test material administration (24-hour reading) and one and two days later (48-hour and 72-hour reading). Scoring of irritative effects was performed according to the method of Draize (1959), in which corneal, iris, and conjunctival effects are scored separately. An irritation score was calculated for each rabbit on a basis of 0-110. Results indicate that the eyes of none of the six rabbits were found to show evidence of positive corneal, iris, or conjunctival changes. Irritation scores in individual rabbits were all zeros. In conclusion, the material is not classified as an irritant following ocular application.

#### **5.7.2 Guinea Pig Maximization Test (Magnusson and Kligman Method) (Hill Top Biolabs Report No. 87-1064-21 (B); E-059)**

This study was conducted to evaluate the potential of EPG-08-oleate (lot # 606617 THPG08), to cause delayed contact hypersensitivity in guinea pigs (Hartley albino) using the methodology of Magnusson and Kligman. The test material is a clear, light yellow liquid. Twenty test animals (weighing 319 to 871 g), ten positive control animals, and ten vehicle control animals were injected intradermally on day 0 with preparations of test material, formaldehyde, and acetone, respectively. Each animal also received two intradermal injections of 50% v/v Freund's Complete Adjuvant (FCA) in distilled water and two injections of the respective test or control materials in the FCA/distilled water emulsion. On day 6, these same groups of animals were exposed topically to a preparation of their respective test or control material. Patches were secured under ELASTOPLAST bandage wrappings for approximately 48 hours. On day 19, all animals were topically challenged at a naive skin site using a preparation of their respective test or control material under ELASTOPLAST bandage wrappings for approximately 24 hours. Naive control animals were patched identically to and concurrently with the test and positive control animals. On the day after removal, the sites were depilated. Later that day and again the next day, the sites were graded for erythema responses (24- and 48-hour responses). On day 27, all of the original test animals were topically rechallenged at a naive skin site using a preparation of the test material under ELASTOPLAST bandage wrapping for approximately 24 hours. Naive control animals were patched identically to and concurrently with the test animals. On the day after removal, the sites were depilated. Later that day and again the next day, the sites were graded for erythema responses (24- and 48-hour responses).

Following primary challenge, the incidence of grade 1 responses in the test group (0 of 20) was compared to that of the naive test control group (0 of 10). The incidence of these responses was comparable to that produced by the naive control group and resulted in a classification of weak sensitization. However, due to an increased incidence of  $\pm$  reactions (slight patchy erythema) in

the test group, (15 of 20), as compared to the naive test control group, (3 of 10), a rechallenge was performed to more clearly define the mechanism. Following primary challenge, the incidence of grade 1 response or greater in the positive control group (10 of 10) was compared to that of the naive positive control group (0 of 10). The incidence of these responses was more pronounced than that produced by the naive positive control group and resulted in a classification of extreme sensitization. Following primary challenge, the incidence of grade 1 responses in the vehicle control group (0 of 10) resulted in a classification of weak sensitization. Eight days following the primary challenge application the twenty test animals were single patch rechallenged. Ten naive control animals were patched identically to and concurrently with the test animals. On the day after removal, the sites were depilated. Later that day and again the next day, the sites were graded for erythema responses (24- and 48-hour responses).

Following the single patch rechallenge, the incidence of grade 1 responses in the test group (0 of 20) was compared to that of the naive control group (0 of 10). The incidence of these responses was comparable to that produced by the naive control group. The incidence of  $\pm$  reactions in the test group, (16 of 20), as compared to the naive test control group, (3 of 10), remained essentially unchanged from that of primary challenge confirming a classification of weak sensitization. As indicated above, classification in accordance with the protocol categorizes the test and vehicle materials as those exhibiting a weak rate of sensitization. It is important to note that this category includes materials which have induced a 0% sensitization rate. This 0% sensitization rate is consistent with the activity of EPG-08-oleate (lot # 606617 THPG08), and the acetone vehicle under the conditions of this protocol. In conclusion, under the conditions of this study, there is no evidence to suggest that EPG-08-oleate (lot # 606617 THPG08) is a dermal sensitizer.

## **5.8 Genotoxicity Testing of Esterified Propoxylated Glycerol (EPG) (Bechtel, 2014)**

### **5.8.1 Bacterial reverse mutation assays (*S. typhimurium* and *E. coli*)**

#### **5.8.1.1 Study to determine the ability of H-EPG-05 HR/SO (9:1) to Induce Mutation in Four Histidine-Requiring Strains of *Salmonella Typhimurium* and Two Tryptophan-Requiring Strains of *Escherichia Coli* (CLE study number ACU 1/S; E-026)**

The ability of H-EPG-05 HR/SO (9:1) to induce mutations in 4 histidine-requiring strains (TA98, TA100, TA1535, and TA1537) of *Salmonella typhimurium* and 2 tryptophan-requiring strains (WP2 pKM101 and WP2 uvrA Pkm101) of *Escherichia coli* (*E. Coli*) was tested in the presence and absence of metabolic activation from the rat liver post-mitochondrial fraction (S9) in two separate Ames assays. Negative (acetone) and positive controls [2-nitrofluorene (2NF), sodium

azide (NaN<sub>3</sub>), 9-aminoacridine (AAC), 4-nitroquinoline 1-oxide (NQO), 2-aminoanthracene (AAN), depending on the strain of the bacteria] were also used.

Based on results of a toxicity range-finder experiment conducted in TA100 (results not discussed herein), *Salmonella* and *E. Coli* strains were exposed to 1.6, 8, 40, 200, or 1000 µg/plate of H-EPG-05 HR/SO (9:1), with and without S9, in Experiment 1. Precipitation of EPG was observed at the highest concentration. In addition, a slight thinning of the bacterial lawn in strain TA98 when exposed to 1000 µg/plate of H-EPG-05 HR/SO (9:1) in the absence of S9, was considered a toxic effect. In Experiment 2, *Salmonella* and *E. Coli* strains were exposed to 62.5, 125, 250, 500, or 1000 µg/plate of H-EPG-05 HR/SO (9:1) with and without S9; treatments in the presence of S9 included a pre-incubation step prior to plating. Precipitation of H-EPG-05 HR/SO (9:1) was observed at the 1000 µg/plate concentration level in all strains, as well as at the 500 µg/plate concentration level in *E. Coli* strain WP2 *uvrA* pKM101 only. No evidence of toxicity was observed.

The number of revertant colonies in the negative control treatments fell within normal ranges and the number of revertant colonies in the positive control treatments increased dramatically. There were no reproducible increases in revertant colonies in the presence and absence of S9 in *Salmonella* and *E. Coli* strains following treatment with H-EPG-05 HR/SO (9:1) at concentrations up to its limit of solubility. As such it was concluded under the conditions of this study H-EPG-05 HR/SO (9:1) is not mutagenic.

#### 5.8.1.2 Additional Studies

Separate Ames assays were conducted using the same protocol to evaluate the mutagenicity of H-EPG-05 soyate<sup>7</sup> and H-EPG-14 soyate<sup>8</sup>. Bacterial strains were exposed to 1.6, 8.0, 40, 200 and 1,000 µg/plate (Experiment 1) and 62.5, 125, 250, 500 and 1,000 µg/plate (Experiment 2) of each form of EPG in the presence and absence of metabolic activation. Precipitation of EPG was restricted to the highest concentration and no evidence of mutagenicity was observed in assays conducted with either H-EPG-05 soyate or H-EPG-14 soyate.

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<sup>7</sup> Study to determine the ability of H-EPG-05 Soyate to Induce Mutation in Four Histidine-Requiring Strains of *Salmonella Typhimurium* and Two Tryptophan-Requiring Strains of *Escherichia Coli*. (CLE study number ACU 2/S)

<sup>8</sup> Study to determine the ability of H-EPG-14 Soyate to Induce Mutation in Four Histidine-Requiring Strains of *Salmonella Typhimurium* and Two Tryptophan-Requiring Strains of *Escherichia Coli*. (CLE study number ACU 3/S).

## 5.8.2 Bacterial reverse mutation assay (*S. typhimurium*)

### 5.8.2.1 *Mutagenicity Test on Heated EPG-05 HR/SO (9:1), Unheated EPG-05 HR/SO (9:1), Heated Palm/Rapeseed Oil (60:40), and Unheated Palm/Rapeseed Oil (60:40) in the Salmonella/Mammalian-Microsome Reverse Mutation Assay (Ames Test) with a Confirmatory Assay*

A modified (preincubation) Ames assay was used to assess the mutagenic potential of H-EPG-05 HR/SO 9:1, heated and unheated vs. palm/rapeseed (60:40), heated and unheated. *S. typhimurium* histidine strains TA98 and TA100 were incubated with the test materials at concentrations of 100, 250, 500, 1000, 2500, and 5000 µg/plate, in the presence and absence of S9 mix. Vehicle (acetone) and positive controls (AAN, 2NF, NaN<sub>3</sub>) were tested concurrently in the presence and absence of S9.

The study consisted of two independent tests, the initial mutagenicity assay and a confirmatory assay (Experiments 1 and 2, respectively). A response was considered positive if the test material produced a concentration-related increase of at least two-fold in the mean number of revertants per plate in at least one of the tester strains over the appropriate vehicle control value.

Heated and unheated versions of H-EPG-05 HR/SO 9:1 did not increase the number of histidine revertants per plate of *S. typhimurium* histidine strains TA98 and TA100 exposed in the presence and absence of S9. On this basis it was concluded that, under the conditions of this assay, heated and unheated versions of H-EPG-05 HR/SO 9:1 were not mutagenic.

### 5.8.2.2 *Additional Assay*

A separate modified (preincubation) Ames assay was conducted using the same protocol to evaluate the mutagenicity of EPG-05 HR/ST 45:55, heated and unheated vs. cottonseed oil, heated and unheated<sup>9</sup>. *S. typhimurium* histidine strains TA98 and TA100 were incubated with the test material at concentrations of 100, 250, 500, 1000, 2500, and 5000 µg/plate, in the presence and absence of S9 mix. Under the conditions of this assay, heated and unheated EPG-05 HR/ST 45:55 were not mutagenic.

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<sup>9</sup> Mutagenicity test on EPG-05 Solid Heated HR/ST (45:55), EPG-05 Solid Unheated HR/ST (45:55), Heated Cottonseed Oil, and Unheated Cottonseed Oil in the Salmonella/Mammalian-Microsome Reverse Mutation Assay (Ames Test) with a Confirmatory Assay Preincubation Method.

### 5.8.3 Mouse Lymphoma Assays

#### 5.8.3.1 *Study to Determine the Ability of H-EPG-05 HR/SO (9:1) to Induce Mutations at the Thymidine Kinase (tk) Locus in Mouse Lymphoma L5178Y Cells Using a Fluctuation Assay (CLE study number: ACU 1/TK; E-017)*

The ability of H-EPG-05 HR/SO (9:1) to induce mutation at the thymidine kinase (tk) locus (5-trifluorothymidine resistance) in L5178Y tk +/- mouse lymphoma cells was evaluated using a fluctuation assay. The assay was performed in the absence and presence of rat liver post-mitochondrial fraction S9. Based on results of a toxicity range-finder experiment (results not discussed) cells were exposed to 0, 12.5, 25, 50, 100, or 200 µg/mL of H-EPG-05 HR/SO (9:1) in Experiment 1 and 2; the highest concentration was selected based on the solubility limit of H-EPG-05 HR/SO (9:1). Negative (acetone) and positive controls [benzo(a)pyrene (with S9) and 4-nitroquinoline 1-oxide (without S9)] were also included. No statistically significant increases in mutants were observed after exposure to H-EPG-05 HR/SO (9:1). Positive control chemicals exhibited statistically significant frequencies of mutants in culture, whereas negative control cultures demonstrated mutants within normal ranges. Thus, it was concluded, that under the conditions of this assay, H-EPG-05 HR/SO (9:1) does not induce mutations, with or without metabolic activation, at the tk locus in mouse lymphoma cells.

#### 5.8.3.2 *Additional Studies*

Separate mouse lymphoma fluctuation assays were conducted using the same protocol to evaluate the mutagenicity of H-EPG-05 soyate<sup>10</sup> and H-EPG-14 soyate<sup>11</sup>. Cells were exposed to H-EPG-05 soyate or H-EPG-14 soyate at concentrations 0, 25, 50, 75 or 100 µg/mL in the presence and absence of S9. No statistically significant increases in mutants were observed after exposure to either form of EPG. Thus, it was concluded, H-EPG-05 soyate and H-EPG-14 soyate do not induce mutations, with or without metabolic activation, at the tk locus in mouse lymphoma cells.

### 5.8.4 Chromosomal Aberrations Assays

#### 5.8.4.1 *Study to Evaluate the Chromosome Damaging Potential of H-EPG-05 HR/SO (9:1) by its Effects on Cultured Human Lymphocytes Using an in Vitro Cytogenetics Assay (CLE Study Number: ACU 1/HLC; E-020)*

An *in vitro* cytogenetics assay using human peripheral blood lymphocyte cultures from a male and a female donor was used to assess the ability of H-EPG-05 HR/SO (9:1) to induce

<sup>10</sup> Study to Determine the Ability of H-EPG-05 Soyate to Induce Mutations at the Thymidine Kinase (tk) Locus in Mouse Lymphoma L5178Y Cells Using a Fluctuation Assay (CLE study number: ACU 2/TK)

<sup>11</sup> Study to Determine the Ability of H-EPG-14 Soyate to Induce Mutations at the Thymidine Kinase (tk) Locus in Mouse Lymphoma L5178Y Cells Using a Fluctuation Assay (CLE study number: ACU 3/TK)

structural aberrations in two separate experiments. Both experiments were performed in the absence and presence of rat liver post-mitochondrial fraction S9. Negative (acetone) and positive controls [4-nitroquinoline 1-oxide (without S9) and cyclophosphamide (with S9), following a 20+0 hour exposure and a 3+17 hour exposure, respectively] were also used.

A toxicity range-finder experiment (results not discussed) was conducted in order to establish the concentration range for Experiments 1 and 2; the highest concentration was selected based on the solubility limit of EPG-05 HR/SO (9:1). In experiment 1, cells were exposed to 0.9887 to 50.00 µg/mL of EPG-05 HR/SO (9:1) for 20 hours in the absence of metabolic activation, and 3 hours in the presence of metabolic activation, followed by 17 hours of recovery. Based on the effects of H-EPG-05 HR/SO (9:1) on the mitotic index, cells exposed to 24.5, 35, and 50 µg/mL of EPG-05 HR/SO (9:1) were analyzed to assess the frequency of chromosomal aberrations. Results demonstrated no clear indications of mitotic inhibition.

In Experiment 2, cells were exposed to 2.112 to 50.00 µg/mL of EPG-05 HR/SO (9:1) for 20 hours in the absence of metabolic activation and 3 hours in the presence of metabolic activation, followed by 17 hours of recovery. Due to the indefinite results in Experiment 1, Experiment 2 also included delayed sampling times (44+0 and 3+41) and a pulse treatment (3+17) at the highest concentration. Based on the effects of EPG-05 HR/SO (9:1) on the mitotic index, cells exposed to 28.13, 37.5, and 50 µg/mL of EPG-05 HR/SO (9:1) were analyzed to assess the frequency of chromosomal aberrations. Afterwards, cells were exposed to the highest concentration of 50 µg/mL of EPG-05 HR/SO (9:1) in a delayed harvest study conducted in the presence (3 hours of exposure and 41 hours of recovery) and absence (44 hours of exposure) of metabolic activation, and in a pulse treatment (3 hours of exposure and 17 hours of recovery) study without metabolic activation.

Positive controls demonstrated statistically significant increases in the proportion of cells with structural aberrations, and the negative control demonstrated a proportion of cells with structural aberrations within normal ranges, confirming the validity of the assay. No significant differences were observed either in the absence or presence of metabolic activation between the H-EPG-05 HR/SO (9:1)-treated and negative control cultures. Thus, it was concluded that H-EPG-05 HR/SO (9:1) does not induce chromosome aberrations in cultured human peripheral blood lymphocytes when tested to its limit of solubility.

#### 5.8.4.2 *Additional Studies*

Separate chromosome aberration tests were conducted using the same protocol to evaluate the mutagenicity of H-EPG-05 soyate<sup>12</sup> and H-EPG-14 soyate<sup>13</sup>. Cultures were treated with H-EPG-

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<sup>12</sup> Study to Evaluate the Chromosome Damaging Potential of H-EPG-05 Soyate by its Effects on Cultured Human Lymphocytes Using an *in Vitro* Cytogenetics Assay (CLE Study Number: ACU 2/HLC)

05 soyate, at concentrations ranging from 0.9887 to 50.00 µg/mL (Experiment 1) and 6.674 to 50.00 µg/mL (Experiment 2). Cells exposed to 24.5, 35, and 50 µg/mL of H-EPG-05 soyate in Experiment 1 and 28.13, 37.5, and 50 µg/mL of H-EPG-05 soyate in Experiment 2 were analyzed to assess the frequency of chromosome aberrations. A small, but statistically significant increase in aberrant cells was observed in Experiment 1 following treatment with H-EPG-05 soyate, however, it was not considered biologically significant because it was not reproducible in Experiment 2. Due to the indefinite results in Experiment 1, Experiment 2 also included delayed sampling times (44+0 and 3+41) and a pulse treatment (3+17) at the highest concentration of 50.00 µg/mL. With respect to H-EPG-14 soyate, cultures were treated with the test material at concentrations ranging from 1.186 to 60.00 µg/mL (Experiment 1) and 8.009 to 60.00 µg/mL (Experiment 2), in the presence and absence of metabolic activation. Cells exposed to 29.4, 42, and 60 µg/mL of H-EPG-14 soyate in Experiment 1 and 29.4, 42, and 60 µg/mL of H-EPG-14 soyate in Experiment 2 were analyzed to assess the frequency of chromosome aberrations. Due to the indefinite results in Experiment 1, Experiment 2 also included delayed sampling times (44+0 and 3+41) and a pulse treatment (3+17) at the highest concentration of 60.00 µg/mL. A small, but statistically significant increase in aberrant cells was observed in Experiment 2; however, it was not considered biologically significant since the number of aberrant cells present were within normal range of traditional values for the negative control. Thus, it was concluded that H-EPG-05 soyate and H-EPG-14 soyate did not induce chromosome aberrations in cultured human peripheral blood lymphocytes when tested to their limit of solubility.

### 5.8.5 Unscheduled DNA Synthesis Assays

#### 5.8.5.1 *Study to Evaluate the Potential of H-EPG-05 HR/SO (9:1) to Induce Unscheduled DNA Synthesis in Rat Liver Using an in Vivo/in Vitro Procedure (CLE study number ACU 1/ILU; E-023)*

This study investigated the ability of H-EPG-05 HR/SO (9:1) to induce unscheduled DNA synthesis (UDS) in the livers of rats following oral administration. Groups of six male Wistar rats (46 to 59 days old, weighing 214 to 329 g) were treated with the negative control, positive control, or 632.5 and 2000 mg/kg of H-EPG-05 HR/SO (9:1), *via* oral gavage, in two experiments. Corn oil, the vehicle for H-EPG-05 HR/SO (9:1), was used as the negative control chemical; positive controls were 2-acetamidofluorene (75 mg/kg, 12-14 hour exposure) and dimethylnitrosamine (10 mg/kg, 2-4 hour exposure) for Experiments 1 and 2, respectively. A toxicity range-finder experiment (results not discussed) was performed to obtain concentration levels for Experiments 1 and 2. In both Experiment 1 and 2, animals were sacrificed and their livers were used to establish a primary hepatocyte culture. Cultures were treated with [<sup>3</sup>H]

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<sup>13</sup> Study to Evaluate the Chromosome Damaging Potential of H-EPG-14 Soyate by its Effects on Cultured Human Lymphocytes Using an *in Vitro* Cytogenetics Assay (CLE Study Number: ACU 3/HLC)

thymidine, slides were fixed with hepatocytes, and dipped with photographic emulsion to prepare autoradiograms. Slides were examined microscopically to calculate the net grains/nucleus (NG = nuclear grain count – mean cytoplasmic grain count) and percentage of cells in repair (net grain  $\geq$ 5) for each slide, animal, and concentration group.

Negative control animals demonstrated a mean NG value less than 0 with only 0 to 0.6% cells in repair. The positive control animals demonstrated mean NG values greater than 5 with more than 50% cells in repair. Treatment with 632.5 or 2000 mg/kg of H-EPG-05 HR/SO (9:1) produced mean NG values no more than -0.9 with no more than 1.4% cells in repair. On this basis, it was concluded that H-EPG-05 HR/SO (9:1) was not genotoxic under the conditions of this assay.

#### 5.8.5.2 Additional Studies

Separate *in vivo/in vitro* unscheduled DNA synthesis tests were conducted using the same protocol to evaluate the mutagenicity of H-EPG-05 soyate<sup>14</sup> and H-EPG-14 soyate<sup>15</sup>. Negative control animals demonstrated a mean NG value less than 0 with only 0 to 1.8% cells in repair in both studies. Treatment with 632.5 or 2000 mg/kg of H-EPG-05 soyate or H-EPG-14 soyate produced mean NG values no more than -0.5 with no more than 0.4% cells in repair, and mean NG values no more than -0.8 with no more than 1.4% cells in repair, respectively. Therefore, it was concluded that H-EPG-05 soyate and H-EPG-14 soyate were not genotoxic under the conditions of these assays.

## 6.0 HUMAN SAFETY

Studies of human tolerance to EPG (H-EPG-05 HR/SO 9:1 and EPG-05 HR/ST 45:55), including single dietary exposure studies and incremental increasing multiple dietary exposure studies, demonstrated that food products prepared with EPGs were highly palatable compared to similar foods prepared with conventional fats. Furthermore, no untoward effects in human volunteers resulted from the consumption of up to 150 g of EPG per day.

It is worthwhile to note that a decline in serum HDL was observed in some subjects placed on very high EPG diets, which greatly reduced the available fat in their diet. This effect was consistent with reports in the published scientific literature which describe a transient drop in serum HDL in subjects placed on low fat diets (Knuiman *et al.*, 1987; Schaefer *et al.*, 1981; Brinton *et al.*, 1990).

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<sup>14</sup> Study to Evaluate the Potential of H-EPG-05 Soyate to Induce Unscheduled DNA Synthesis in Rat Liver Using an *in Vivo/in Vitro* Procedure (CLE study number ACU 2/ILU)

<sup>15</sup> Study to Evaluate the Potential of H-EPG-14 Soyate to Induce Unscheduled DNA Synthesis in Rat Liver Using an *in Vivo/in Vitro* Procedure (CLE study number ACU 3/ILU)



Table 6-1 provides a summary of human studies conducted with EPG; individual study summaries are provided in Sections 6.1 and 6.2. Articles based on the studies entitled, "Assessment of the Effect of Esterified Propoxylated Glycerol (EPG) on the Status of Fat-Soluble Vitamins and Select Water-soluble Nutrients following Dietary Administration to Humans for Eight Weeks" (Section 6.2.3), and "Tolerance of Rising Dietary Concentrations of Esterified Propoxylated Glycerol (EPG) Among Human Volunteers" (Section 6.1.1), have been published in *Regulatory Toxicology and Pharmacology*.

**Table 6-1 Summary of human studies with EPG in foods**

Study ID	Type of exposure	EPG consumption	Number of subjects completed the study	Observations
ICR 0047047*	Rising EPG intake (with 2-day wash-out in between) H-EPG-05 HR/SO 9:1	Day 4: 0 g Day 7: 30 g Day 10: 60 g Day 13: 90 g Day 16: 120 g Day 19: 150 g	16	No effects on vital signs, hematology, or urinalysis.  EPG was associated with ↓HDL cholesterol in all subjects; transient ↑liver transaminase (12 subjects).  Except for flatus, GI-related adverse events decreased with decreasing fat and increasing EPG concentrations.
ICR 004251	Constant EPG intake H-EPG-05 HR/SO 9:1	Days 1-18: 120 g/day	15	No effects on hematology or urinalysis.  EPG was associated with ↓HDL cholesterol in all subjects; transient ↑liver transaminase (10 subjects).  Greater frequency of bowel movements after Day 4 (all subjects), abdominal pain (11 subjects), and oil leakage from rectum (3 subjects).  120 g/day considered well tolerated.
ICR 010197	Constant EPG intake EPG-05 HR/ST 45:55	Days 1-3: 0 g/day Days 4-21: 60 g/day	12	No effects on vital signs, hematology, urinalysis, body weight, serum 25-OH vitamin D, or ECG.
	Declining EPG intake EPG-05 HR/ST 45:55	Days 1-3: 0 g/day Days 4-7: 120 g/day Days 8-16: 100 g/day Days 17-21: 76 g/day	14	EPG was associated with ↓HDL cholesterol in 7 subjects in Group 1 and 6 subjects of Group 2; ↑liver transaminase (AST, ALT).  Three subjects withdrawn (1 receiving 60 g/day; 2 receiving ≤120 g/day) due to GI adverse events (abdominal pain, nausea, etc.) possibly/probably related to EPG. The most common GI-related adverse event was bowel movement with oil and colored matter. Greater frequency of bowel movements and incidence of abnormal bowel movements as study progressed.  60 g/day considered well tolerated.

\* Bechtel, 2015

**Table 6-1 Summary of human studies with EPG in foods (cont'd)**

Study ID	Type of exposure	EPG consumption	Number of subjects completed the study	Observations
ICR 011426	Constant EPG intake compared to margarine (ordinary triglycerides)  EPG-05 HR/ST 45:55	Days 1-5: 0 g/day Days 6-23: 0, 10, 20, 30, or 40 g/day	10-12 per group	No effects on vital signs, clinical chemistry, hematology, or urinalysis.  ↑Mean body weights in subjects receiving either margarine or 10 g/day EPG; ↓mean body weight in remaining groups.  Statistically analysis showed no significant difference in the frequency of gastrointestinal symptoms for any EPG group or relationship to concentration (↑incidence of difficulty swallowing, excessive flatus, and vomiting at 10 and 40 g/day; ↑vomiting at 30 g/day).  EPG was considered well tolerated.
CCRC EC-10*	Constant EPG intake compared to margarine (ordinary triglycerides)  EPG-05 HR/ST 45:55	Days 1-56: 0, 10, 25, or 40 g/day	34-36 per group	No effects on vital signs, body weight, hematology, serum chemistry, urinalysis, PT, PTT, or circulating retinol, α-tocopherol, folate, vitamin B <sub>12</sub> , zinc, iron, calcium, phosphorus, osteocalcin, RBP, and PTH.  EPG was associated with ↓β-carotene, ↓phyloquinone, and ↑PIVKA-II; ↑25-OH D <sub>3</sub> but to a lesser extent than margarine.  Greater incidence of GI adverse events (gas with discharge, diarrhea, oily spotting/evacuation/stool, liquid/soft stool) at 25 and 40 g/day.  10 g/day considered well tolerated.

\* Davidson and Bechtel, 2014

## **6.1 Tolerance Studies with H-EPG-05 HR/SO (9:1)**

### **6.1.1 A Rising Multiple Dose Tolerance Dietary Study of a Solid Esterified Propoxylated Glycerol Version H-EPG-05 HR/SO 9:1 in Normal Volunteers (Bechtel, 2015)**

This single-center, domiciled, single-blind, increasing-concentration study was designed to evaluate human tolerance of foods containing a solid form of the fat substitute EPG, H-EPG-05 HR/SO 9:1. Sixteen healthy male volunteers received 0, 30, 60, 90, 120, and 150 g of EPG in baked foods and a butter-like spread, as a rising concentration given over a period of 19 days, each separated by a 2-day wash-out period. Vital signs, blood chemistry, hematology, urinalysis, bowel habit, and adverse events were monitored for 23 days. Corresponding fat levels in the diet for these days were approximately 302 g, 270 g, 239 g, 208 g, 183 g, and 149 g, respectively. EPG replaced 0, 10, 20, 30, 40, and 50% fat in the diet, respectively.

All subjects demonstrated decreased HDL cholesterol concentrations. Twelve out of 16 subjects exhibited increased transaminase level at some point during the study; the levels returned to normal by the end of the study or by the post-study follow-up visit. The authors indicated that adverse events related to gastrointestinal dysfunction were associated with large quantities of food and fat consumed. Adverse events, with the exception of flatus, decreased as the fat content decreased and the amount of EPG increased. No serious adverse events were reported. Overall, EPG was well tolerated.

### **6.1.2 A Repeated Dose, Tolerance Study of Dietary Esterified Propoxylated Glycerol (H-EPG-05 HR/SO 9:1) in Normal Volunteers (ICR Project No. 004251; E-030)**

Fifteen healthy male volunteers received 120 g/day of a solid form of EPG, H-EPG-05 HR/SO 9:1, in a single-center, domiciled, single-blind, repeated, constant concentration, tolerance study. EPG was incorporated into 3 meals and 3 snacks, in foods such as cinnamon buns, chocolate bars, scones, and a butter-like spread administered for 18 days. Hematology, clinical chemistry, and urinalysis were monitored on study Days 7, 14, and 22.

All subjects experienced an increase in the frequency of bowel movements after Day 4, 11 subjects reported abdominal pain at some point, and 3 subjects reported transient oil leakage per rectum. All subjects exhibited a decrease in HDL concentration over time. A transient increase of liver transaminases was observed in 10 subjects, these elevations were attributed to a high carbohydrate intake. Reported adverse events of a gastrointestinal nature demonstrated no trend. As such, it was concluded that ingestion of 120 g per day of EPG for 18 days was not associated with any serious adverse effects. This degree of intake is several fold greater than the intake expected among consumers of EPG-containing food products.

## **6.2 Tolerance Studies with EPG-05 HR/ST 45:55**

### **6.2.1 A Repeated, Parallel Group, Constant Dose Tolerance Study of Two Consumption Levels of Dietary Esterified Propoxylated Glycerol (EPG-05 HR/ST 45:55) AOD2-09 (ICR Project No. 010197; E-035)**

This single-center, domiciled, constant-concentration study assessed the safety and tolerance of a solid form of the fat substitute EPG, EPG-05 HR/ST 45:55, as a dietary component of breakfast, lunch, and dinner snacks, such as cinnamon buns, chocolate bars, scones, and spread. In addition, the study was designed to collect data to support a NOEL for oil leakage per rectum, and for oil separation in bowel movements. Healthy male volunteers were randomized into two parallel groups for 22 days. Group 1 included 12 subjects that received 60 g/day of EPG-05 HR/ST 45:55 on days 4-21 of the study. Group 2 included 16 subjects that received a declining EPG-05 HR/ST 45:55 concentration of 120, 100, and 76 g on days 4-7, 8-16, and 17-21, respectively. Vital signs, blood chemistry, hematology, urinalysis, weight, bowel habit, serum 25-monohydroxy vitamin D, electrocardiogram (ECG) results, and adverse events were monitored.

No serious adverse events were observed. No effects were seen in vital signs, ECG, hematology, and urinalysis. Decreased HDL cholesterol concentrations were observed in 7 subjects from Group 1, and 6 subjects from Group 2. Increases in aspartate aminotransferase (AST) and alanine transaminase (ALT) were observed and could possibly be related to EPG intake. The frequency of bowel movements and incidences of abnormal bowel movements increased as the study progressed. A total of 895 out of 947 adverse events reported were related to the gastrointestinal tract; of these, the most common event was a bowel movement with oil and colored matter. One subject receiving 60 g/day was withdrawn due to abdominal pain. Two subjects from group 2 (receiving 120 g/day initially and then reduced to 100 and 76 g/day) were withdrawn due to rectal fissure bleeding, and abdominal pain/nausea. These effects were considered possibly or probably related to EPG. No NOEL for oil leakage per rectum and for oil separation in bowel movements was established. The authors concluded that ingestion of 60 g/day of EPG-05 HR/ST 45:55 for 18 days was not associated with any serious adverse effects.

### **6.2.2 Protocol Number EC-09/011426. A Double-Blind, Parallel Group, Placebo Controlled Tolerance Study of Four Doses of Dietary Esterified Propoxylated Glycerol (EPG-05 HR/ST 45:55) and Placebo in Normal Volunteers (ICR Project No. 011426; E-038)**

A randomized, domiciled, double-blind, parallel group, placebo-controlled study was conducted to assess the safety and tolerability of EPG-05 HR/ST 45:55 and possibly determine an approximate NOEL. The experimental design included a 5-day run-in period, during which

subjects received ordinary triglycerides, followed by an 18-day double-blind period in which 55 subjects (10 to 12/group) were randomized to receive 0, 10, 20, 30, or 40 g of EPG/day in food products such as spread, chocolate, and muffins. Vital signs, adverse events, gastrointestinal symptoms, bowel movements and stool characteristics, hematology, blood chemistry, and body weight were monitored.

According to the principal investigator, no clinically-significant changes were observed in vital signs, ECGs, urinalysis, or hematology. No increase in abnormal bowel movements or loose soft stools was observed in subjects receiving EPG-05 HR/ST 45:55 when compared to ordinary triglycerides. However, there was an increase in the frequency of normal bowel movements in the group receiving 30 g of EPG-05 HR/ST 45:55/day. There was an increase in mean body weight of subjects receiving triglycerides and 10 g of EPG-05 HR/ST 45:55/day, and a decrease in groups receiving 20, 30, and 40 g of EPG-05 HR/ST 45:55/day. Decreased cholesterol and LDL was observed in groups receiving 30 and 40 g of EPG-05 HR/ST 45:55/day; these decreases were small and not considered clinically significant.

No serious adverse events were reported. Ninety-five minor adverse events were observed, of these, headache was most common. Subjects receiving EPG-05 HR/ST 45:55 also reported difficulty swallowing, excess flatus, and vomiting. One subject receiving 30 g of EPG-05 HR/ST 45:55/day demonstrated increased levels of AST and ALT concentrations on Day 10, possibly related to the test material. Another subject in the same group experienced dyspepsia on Day 9, also possibly related to EPG-05 HR/ST 45:55. No significant concentration-related trend in the occurrence of adverse events was observed. A slight increase in the number of adverse events in the group receiving 40 g of EPG-05 HR/ST 45:55/day was observed; however, this was considered unrelated to the administration of the test material. In conclusion, concentrations of 10, 20, 30, or 40 g of EPG-05 HR/ST 45:55/day were expected to cover the NOEL range. However, a NOEL was not established. Overall, EPG-05 HR/ST 45:55 was considered to be safe and well tolerated.

### **6.2.3 Assessment of the Effect of Esterified Propoxylated Glycerol (EPG) on the Status of Fat-Soluble Vitamins and Select Water-soluble Nutrients following Dietary Administration to Humans for Eight Weeks (Davidson and Bechtel, 2014)**

A double-blind, randomized, controlled study was performed to assess the effect of EPG-05 HR/ST 45:55 on fat-soluble vitamins and select nutrients in human subjects. For 8 weeks, 139 healthy volunteers (34 to 36/group) consumed a core diet providing adequate caloric and nutrient intakes. The diet included items (spread, muffins, cookies, and biscuits) providing EPG (10, 25, and 40 g/day) vs. margarine alone (control).

The variables measured at baseline and regular intervals were: physical exam, including vital signs; body weight; hematology; clinical chemistry; urinalysis; circulating levels of  $\beta$ -carotene, retinol (vitamin A),  $\alpha$ -tocopherol (vitamin E), 25-OH D<sub>2</sub> (vitamin D, ergocalciferol), 25-OH D<sub>3</sub> (vitamin D, cholecalciferol), phyloquinone (vitamin K<sub>1</sub>), PIVKA-II (proteins induced in vitamin K absence), serum folate, RBC (red blood cell) folate, vitamin B<sub>12</sub>, zinc, iron, calcium, phosphorus, osteocalcin, RBP (retinol-binding protein), intact PTH (parathyroid hormone), cholesterol, HDL-C (high-density lipoproteins), LDL-C (low-density lipoproteins), and triglycerides; PT and PTT (partial thromboplastin time); urine zinc, sodium, potassium, creatinine, calcium, and phosphorus; and tolerability. Tolerability was assessed by the incidence of 14 specific gastrointestinal adverse events: passing gas; gas with discharge; abdominal bloat/cramp; heartburn; diarrhea; constipation; urgency of bowel movement; fecal incontinence; oily spotting; oily evacuation; oily stool; liquid stool; soft stool; and hard stool.

Significant declines in  $\beta$ -carotene were seen over time, especially in the EPG groups, but with no apparent relationship to EPG concentration (more severe at 10 g/day and 40 g/day than at 25 g/day). It is possible that the apparent effect of EPG on circulating  $\beta$ -carotene was related to a lower dietary fat intake among subjects receiving EPG, since subjects had difficulty consuming all of the additional fat necessary to fully compensate for what EPG is displaced in the diet. In this case, as a lipid-like material, EPG might have affected the absorption of these nutrients strictly through physicochemical processes, acting as a lipid "sink" during transit in the gastrointestinal tract.

As shown in Figure 6-1, based on the Wilcoxon Rank Sum Test, there were no statistically significant changes from baseline at the primary endpoint (Day 56) in mean retinol levels in evaluable subjects receiving EPG 10, 25, and 40 g/day compared with subjects receiving placebo. Similarly, no other statistically significant differences in the mean change from baseline were noted between the EPG groups and placebo group at Days 14, 28, 42, 56 and the end point analysis with the exception of the EPG 25 g/day group at Day 14 ( $P=0.0141$ ).

Likewise, the Wilcoxon Rank Sum Test revealed significant decreases in mean alpha-tocopherol levels from baseline in the EPG 25 g/day group at Days 14 ( $p=0.498$ ), 28 ( $p=0.0014$ ) and 42 ( $p=0.0001$ ) and in the EPG 40 g/day group at Day 14 ( $p=0.0166$ ), Day 42 ( $p=0.0030$ ) compared to the placebo group. No other statistically significant differences in the change from baseline were noted between the EPG groups and the placebo group at Days 14, 28, 42, and 56. The end point analysis using the last observation carried forward (LOCF) was similar to the Day 56 results for each treatment group (Figure 6-2).

Circulating 25-OH D<sub>3</sub> levels increased over time in the EPG groups, but not to the same degree as the control group, which had an unexpected rise, despite attempts to control endogenous 25-

OH-D<sub>3</sub> synthesis by conducting the study during the winter in Chicago, Illinois, USA (Figures 6-3 and 6-4).

EPG intake was associated with a slight decline in phylloquinone levels across all groups. However, the declines did not exceed 0.1 ng/mL and were not statistically significant within any of the individual groups; statistical significance was observed only when the differences within each EPG group were compared to the differences (none or positive) in the control.

By the end of the study, the levels of circulating proteins induced in vitamin K absence (PIVKA-II) had increased significantly in the EPG 25 and EPG 40 groups, compared to the control; in the EPG 10 group, the difference from baseline was comparable to the difference from baseline in the control. Combined with the phylloquinone results, these data suggest that EPG might have affected the synthesis of vitamin K-dependent clotting factors to some extent, but the changes were small, and there was no indication of any clinical manifestation. The changes in clotting parameters (PT and PTT) from baseline to the end of the study were comparable between the control and EPG groups.

With the exception of transient gastrointestinal discomfort, all adverse events reported by subjects in this study were considered unrelated to EPG. Seven of the 14 pre-defined gastrointestinal adverse events (gas with discharge; diarrhea; oily spotting; oily evacuation; oily stool; liquid stool; soft stool) were reported more frequently by subjects receiving 25 or 40 g/day of EPG, especially females. In general, the incidence and duration of these symptoms correlated with EPG dietary concentration. The results suggest 10 g/day of EPG was reasonably well tolerated.



Figure 6-1 Mean retinol levels over time

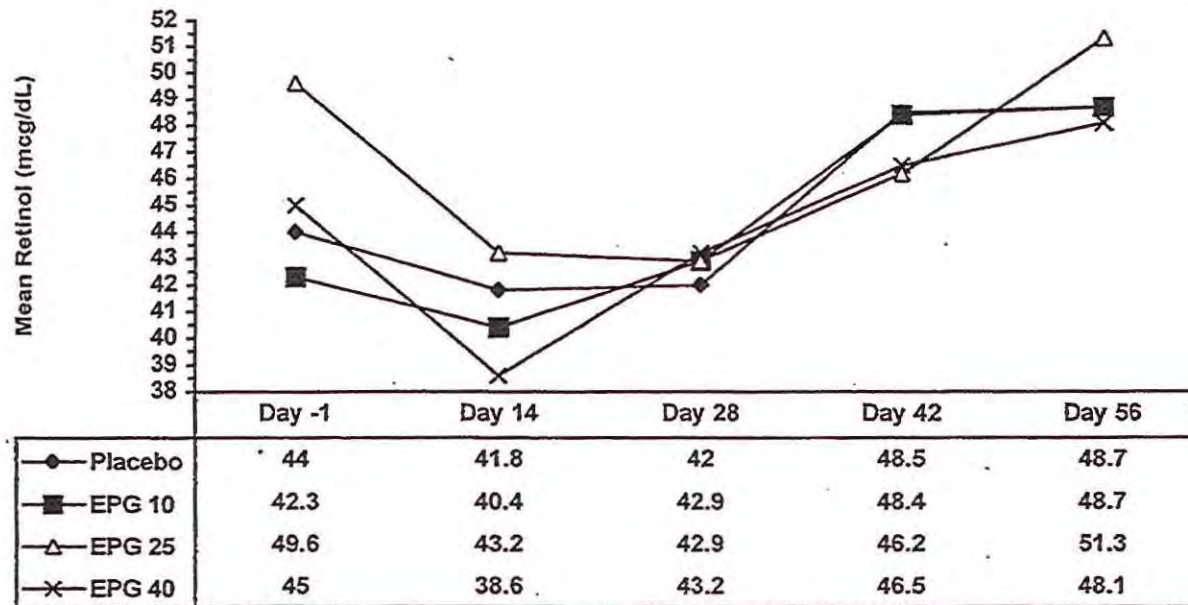
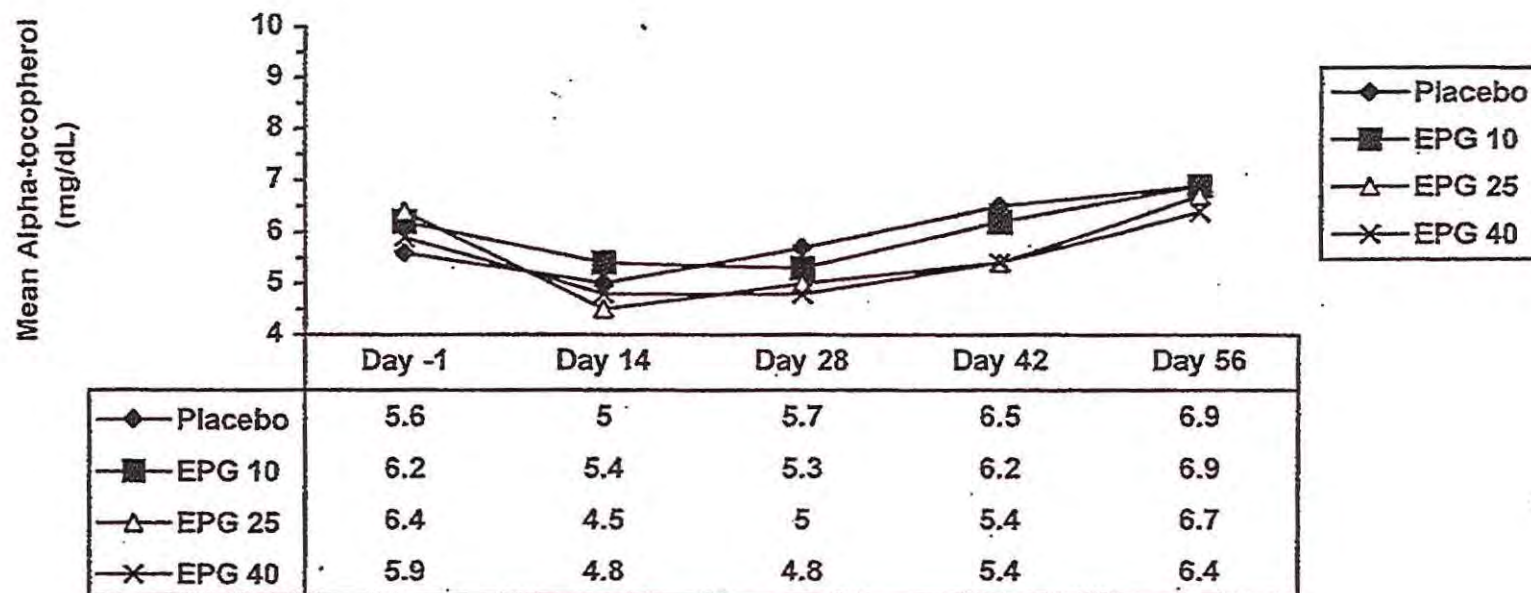
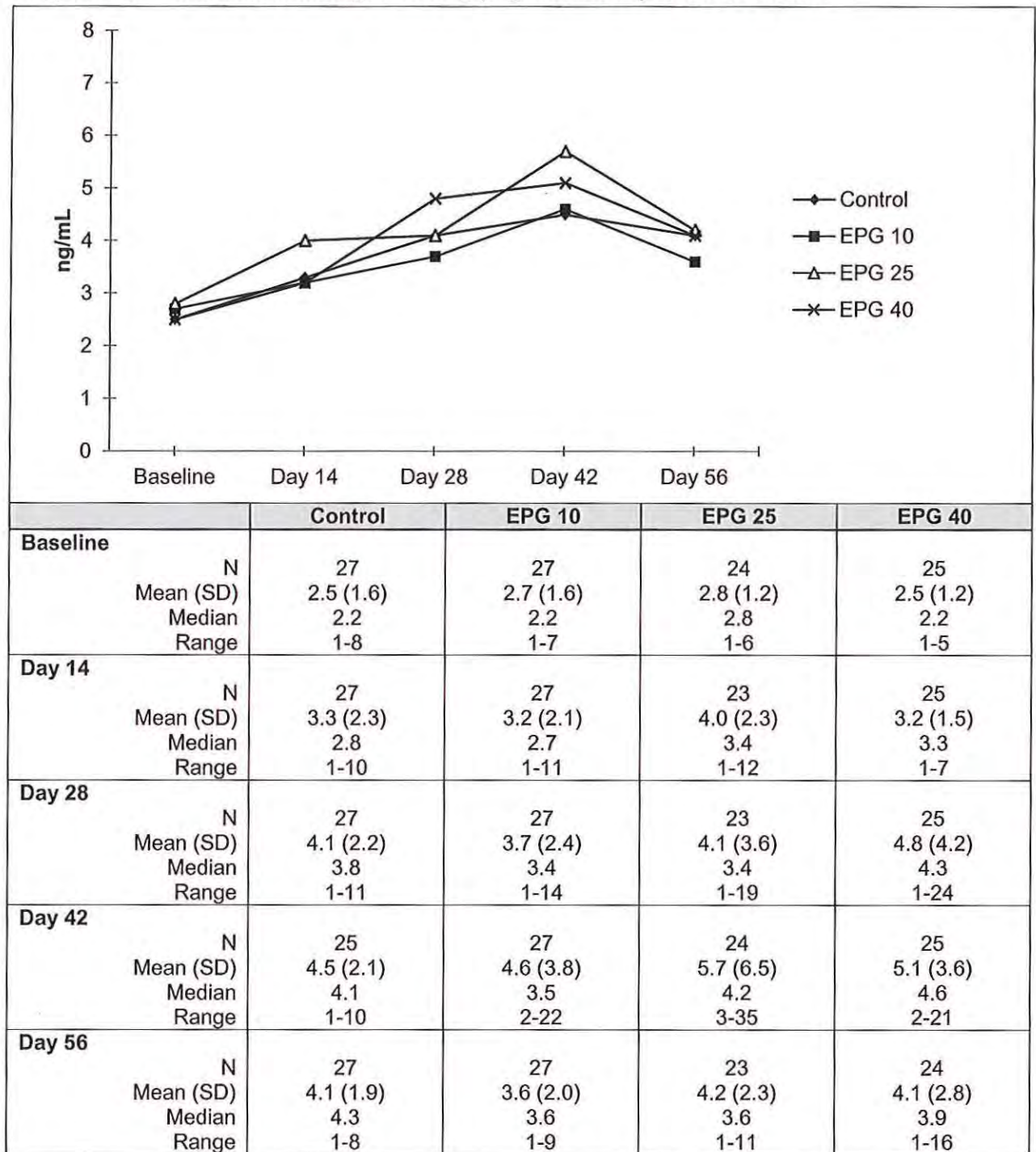


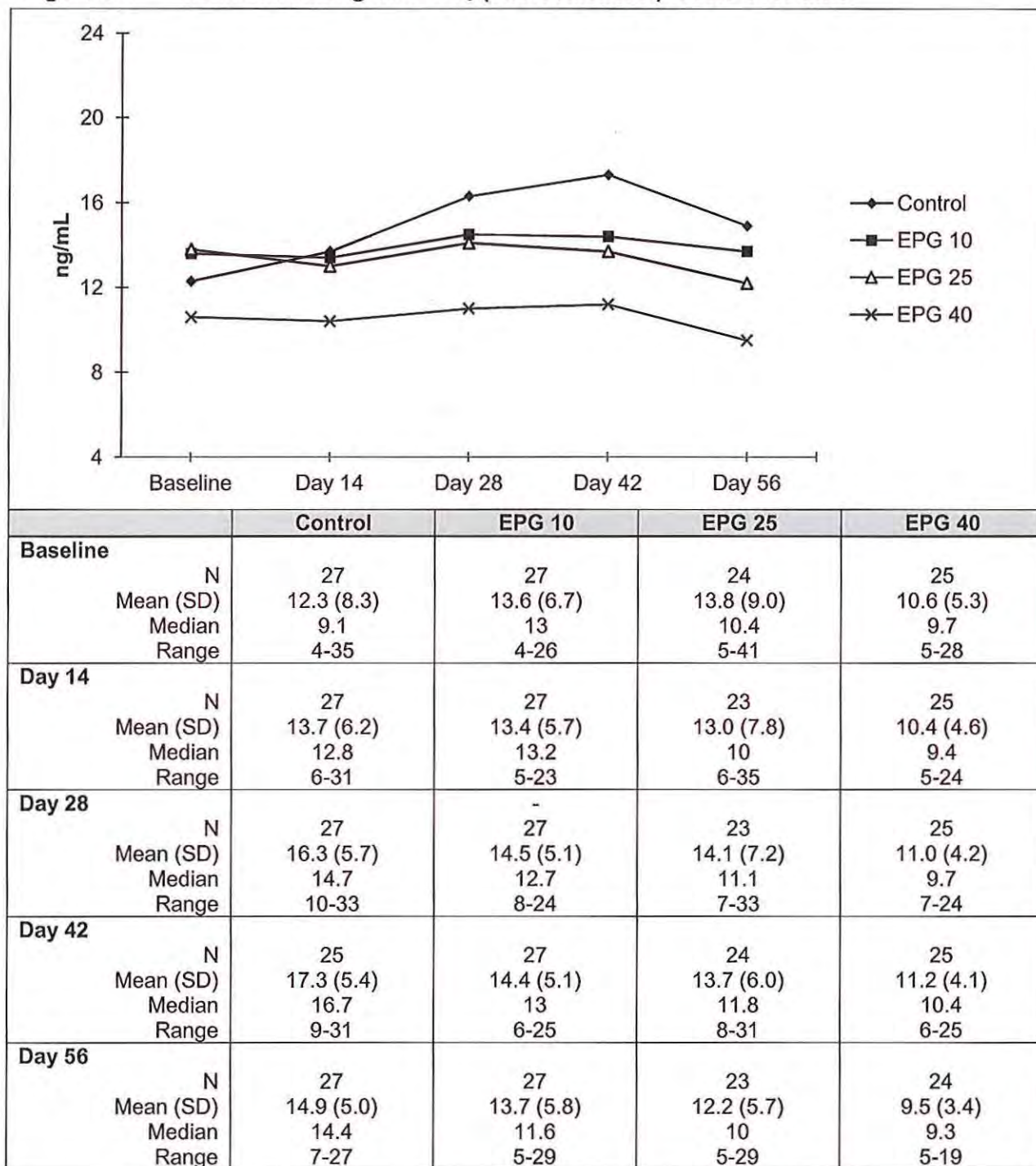
Figure 6-2 Mean alpha-tocopherol levels over time



**Figure 6-3 Mean circulating 25-OH D<sub>2</sub> (ergocalciferol) levels over time**



**Figure 6-4 Mean circulating 25-OH D<sub>3</sub> (cholecalciferol) levels over time**



## **7.0 INFORMATION POTENTIALLY INCONSISTENT WITH GRAS**

### **7.1 Gastrointestinal Discomfort**

Olestra intake has reportedly been associated with gastrointestinal symptoms, including loose stools. This effect is considered similar to the effect of mineral oil, which interferes with the development of firm, well-formed stools (61 FR, 3118; January 30, 1996).

The potential of EPG to induce similar effects was assessed through multiple studies. Administration of solid forms of EPG to experimental animals and humans has not resulted in any adverse effects on gastrointestinal physiology. All versions of EPG currently being considered for use in spreadable and baked goods are solids with melting points at or above 102 °F. As such, the proposed forms are expected to be well tolerated and devoid of adverse gastrointestinal effects.

Occasional separation of the test material from stool bulk has been observed at the highest levels of EPG exposure (up to 150 g/day), but the incidence of loose stool and other gastrointestinal symptoms declines with decreasing dietary concentrations. For example, human volunteers receiving 25 or 40 g/day of a semi-solid form of EPG in food items (spread and baked goods) for eight weeks, reported gastrointestinal adverse events (gas, soft stool, oily spotting, *etc.*) more frequently than subjects receiving margarine alone. However, at 10 g/day, the only statistically significant difference from the control (margarine) group was oily spotting (refer to Sections 6.1 and 6.2).

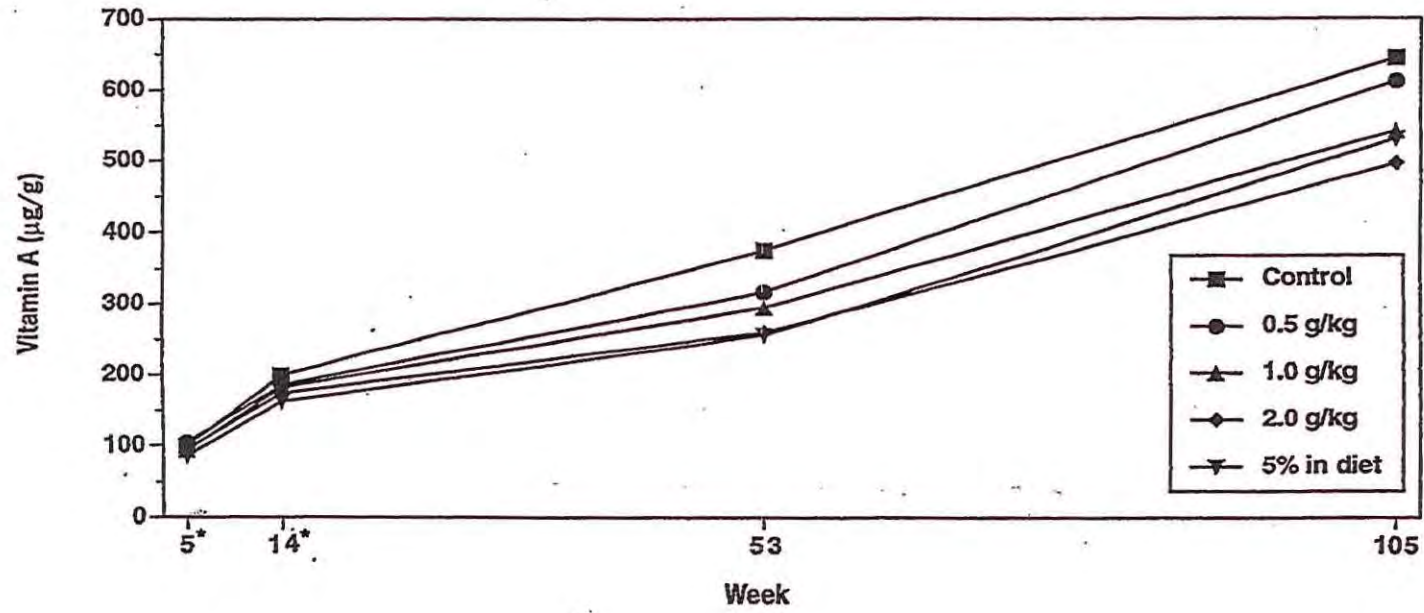
### **7.2 Effect on Nutrient (Fat-Soluble Vitamin) Status**

EPG is intended to replace fats in selected spreadable and baked food products. Some fat-mimetic substances have been shown to interfere with the absorption of lipid-soluble nutrients from the gastrointestinal tract, and conduct of safety studies required the diet to be nutrient fortified. For example, pigs fed olestra for 4 to 26 weeks had lower liver and serum concentrations of vitamins A and E, and lower serum 25-OH vitamin D, in a dose-dependent manner (reviewed by Tulley *et al.*, 2005). Olestra has also been associated with potentially clinically significant lower absorption of fat-soluble vitamins in humans (Schlagheck *et al.*, 1997). Due to the intensity of this nutrient interaction, vitamin fortification of olestra is required by the FDA.

By contrast, EPG exhibits only a weak interaction and none of the safety or clinical studies required diets to be nutrient fortified. Experimental animal studies of EPG have shown some treatment-related effects on the levels of some fat-soluble vitamins (Figures 7-1 through 7-3). Specifically, dietary intake of EPG was associated with lower levels of liver vitamins A (retinol) and E (tocopherol), and serum vitamin D across multiple animal species, in both sexes, and

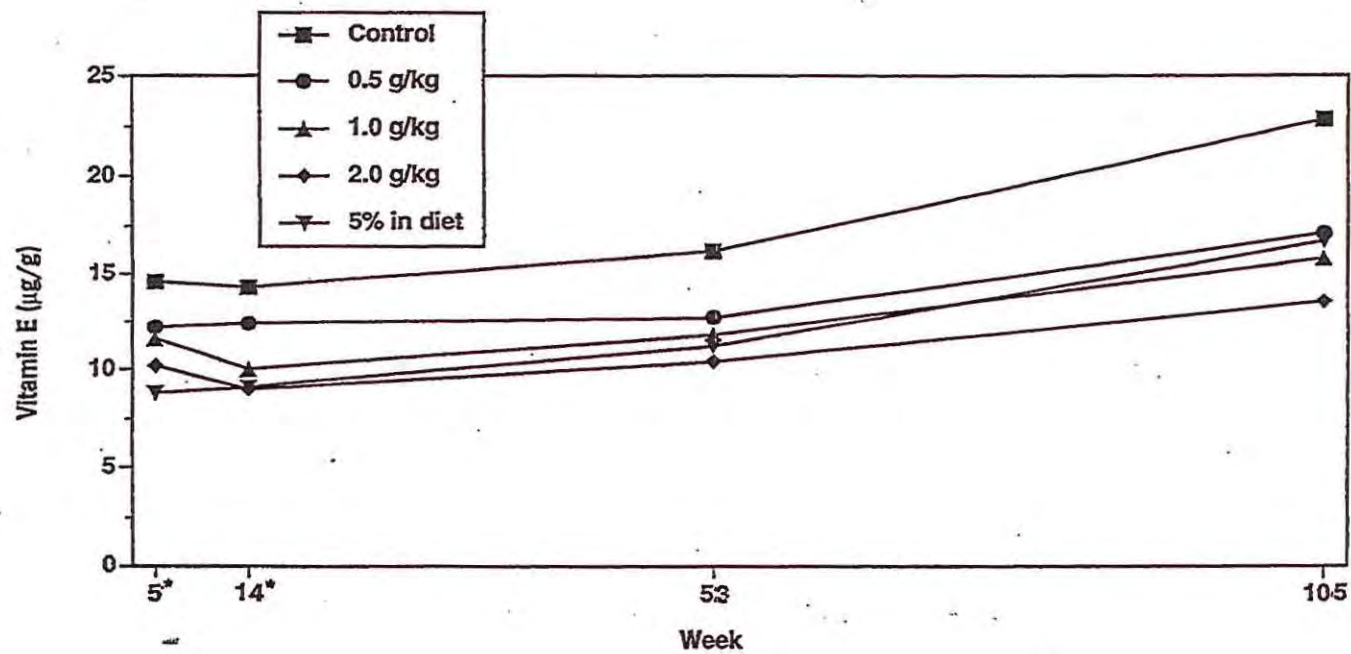
generally in a concentration-dependent manner. Despite these observations, which were statistically significant in many cases, none of the animals in any of the studies exhibited clinical signs or microscopic evidence of vitamin deficiency. By the nature of the study designs, most effects would have been detected. In addition, there was no evidence that the vitamin levels detected in the EPG studies were significantly different from those reported in the published literature for normal control animals. In the long-term EPG studies, the levels of fat-soluble vitamins increased and stabilized over time to fall within normal ranges reported in the literature, despite being significantly lower statistically than those of the corresponding control group animals. As reported in Section 6.2.3, no significant effects were observed in fat soluble vitamin status among human subjects consuming up to 40 g/day of EPG for 8 weeks.

Figure 7-1 Liver vitamin A in male rats



\* Values from HWI 6226-122

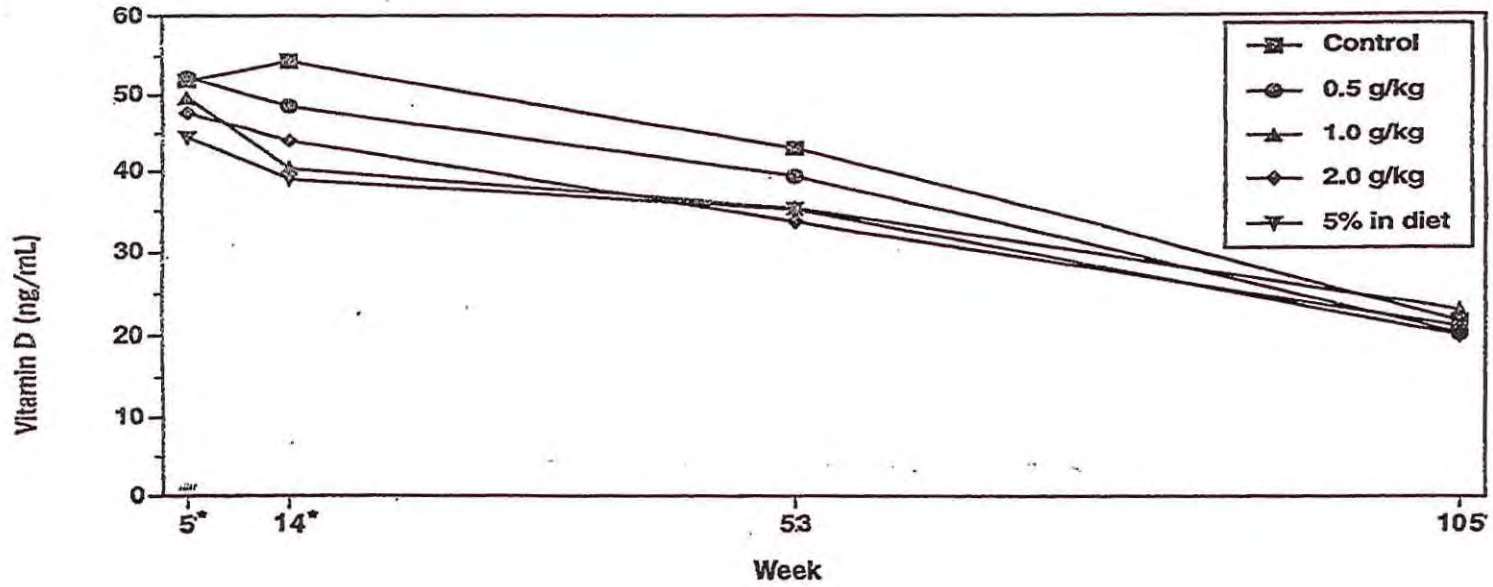
Figure 7-2 Liver vitamin E in male rats



\* Values from HWI 6226-122



Figure 7-3 Serum vitamin D in male rats



\* Values from HWI 6226-122

### 7.3 Effect on $\beta$ -Carotene Levels

As noted in Section 6.2.3, significant declines in  $\beta$ -carotene were reported in subjects receiving EPG; however, if the effect was indeed related to EPG intake, it is uncertain why there was no apparent relationship to EPG concentration (more severe at 10 g/day and 40 g/day than at 25 g/day).

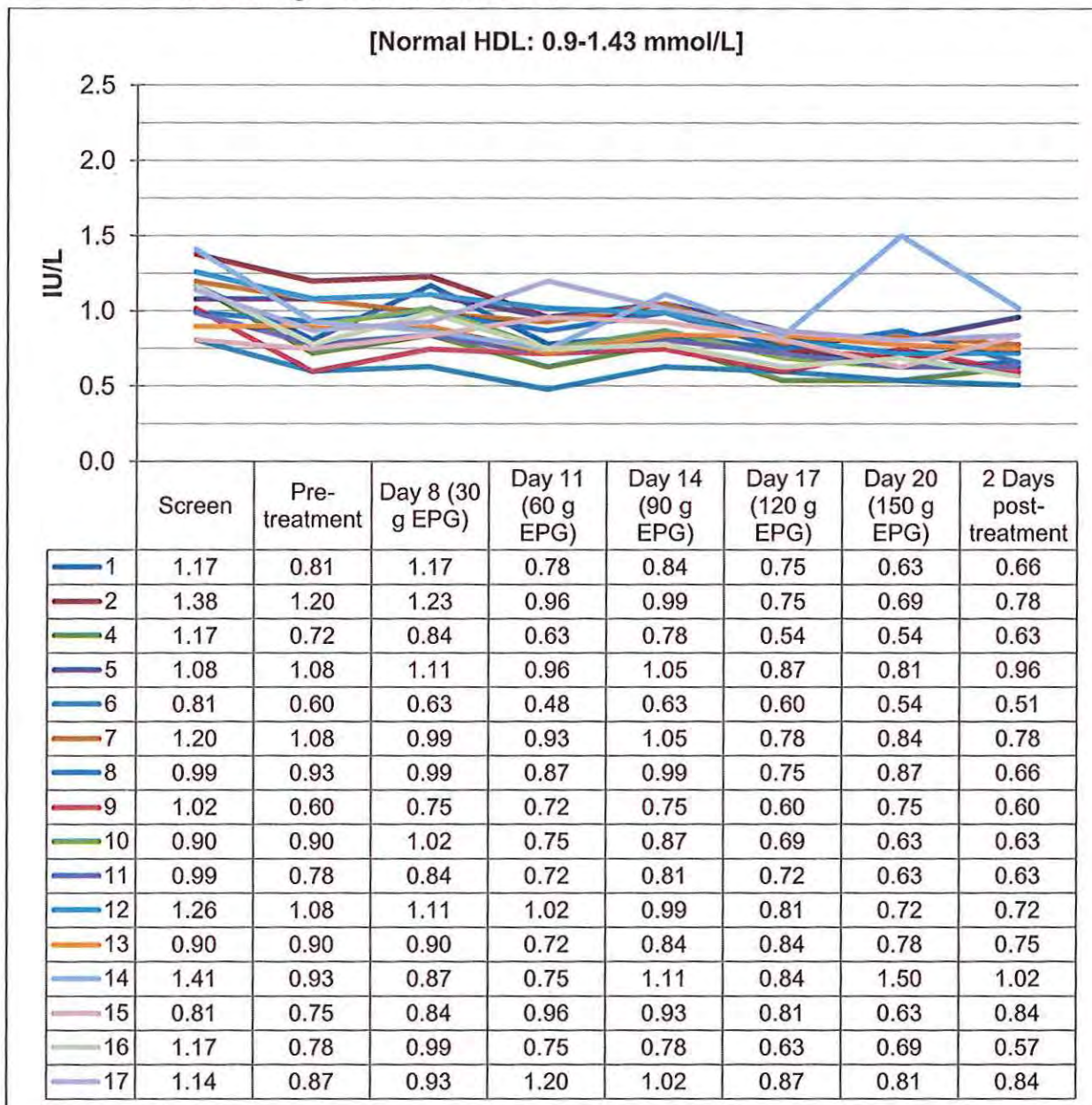
The bioavailability of carotenoids such as  $\beta$ -carotene can vary considerably based on the food matrix, concurrent fat intake, and serum/tissue concentrations. A study that explored, among others, the fate of  $\beta$ -carotene from carrot purée when administered to healthy human subjects, showed based on samples of blood, and stomach and duodenal contents, that the stomach initiates the transfer of carotenoids from the food matrix to the fat phase of a meal, but that the proportion of carotenoids recovered in the micellar phase of the duodenum is very low (<7%) (Tyssandier *et al.*, 2003).

It is possible that the apparent effect of EPG on circulating  $\beta$ -carotene was related to the lower dietary fat intake among subjects receiving EPG. As previously mentioned, subjects might not have consumed all of the additional fat necessary to fully compensate for what EPG displaced in the diet. In this case, as a lipid-like material, EPG might have affected the absorption of these nutrients strictly through physicochemical processes, acting as a lipid “sink” during transit in the gastrointestinal tract. Substances known to reduce the bioavailability of carotenoids are lipid-lowering agents such as cholestyramine and probucol (Elinder *et al.*, 1995), nonabsorbable fat substitutes such as sucrose polyester (olestra) (Peters *et al.*, 1997; Schlagheck *et al.*, 1997; Tulley *et al.*, 2005; Neuhouser *et al.*, 2006), plant sterol-enriched margarines (Gylling *et al.*, 1999; Law, 2000; Hendriks *et al.*, 2003), and dietary fiber supplementation (Rock and Swendseid, 1992). No dietary reference intakes (DRIs) *per se* have been proposed by the Institute of Medicine for carotenoids, although existing recommendations for increased consumption of carotenoid-rich fruits and vegetables are supported (IOM, 2000).

### 7.4 Effect on Serum HDL Levels

Serum HDL levels below the normal range were reported in some individuals receiving EPG in amounts between 60 and 150 g EPG/day for up to 18 days (Figure 8-4). The decrease in serum HDL levels was small and not reported in subsequent studies at lower doses for more extended periods. This effect is consistent with reports in the published scientific literature which describe a transient drop in serum HDL in subjects placed on low fat diets (Knuiman *et al.*, 1987; Schaefer *et al.*, 1981; Brinton *et al.*, 1990). In addition, studies with EPG in micropigs, a species considered to be a good model for human digestion, for up to 1 year did not produce evidence of an effect on serum HDL (Figures 7-5a and 7-5b).

**Figure 7-4 HDL cholesterol levels over time in each of 16 subjects receiving increasing EPG concentrations**



\* Values from ICR Project No. 0047047 (Section 6.1.1)

Figure 7-5a Mean HDL values over time in male micropigs

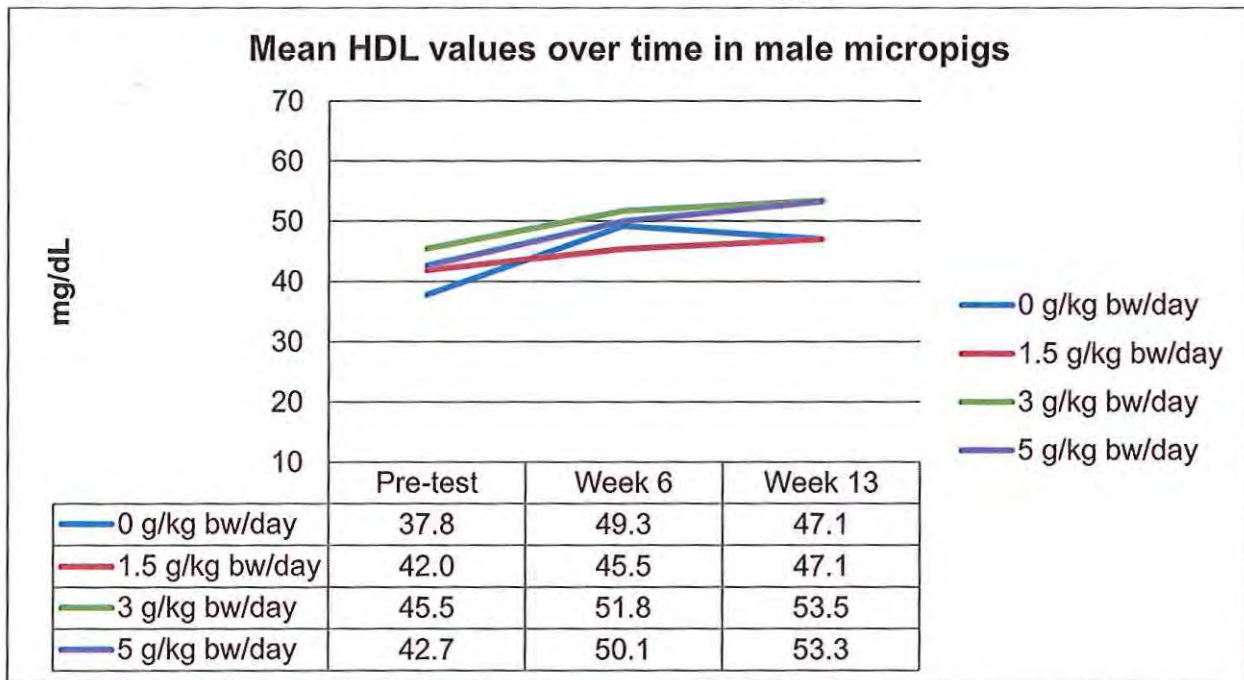
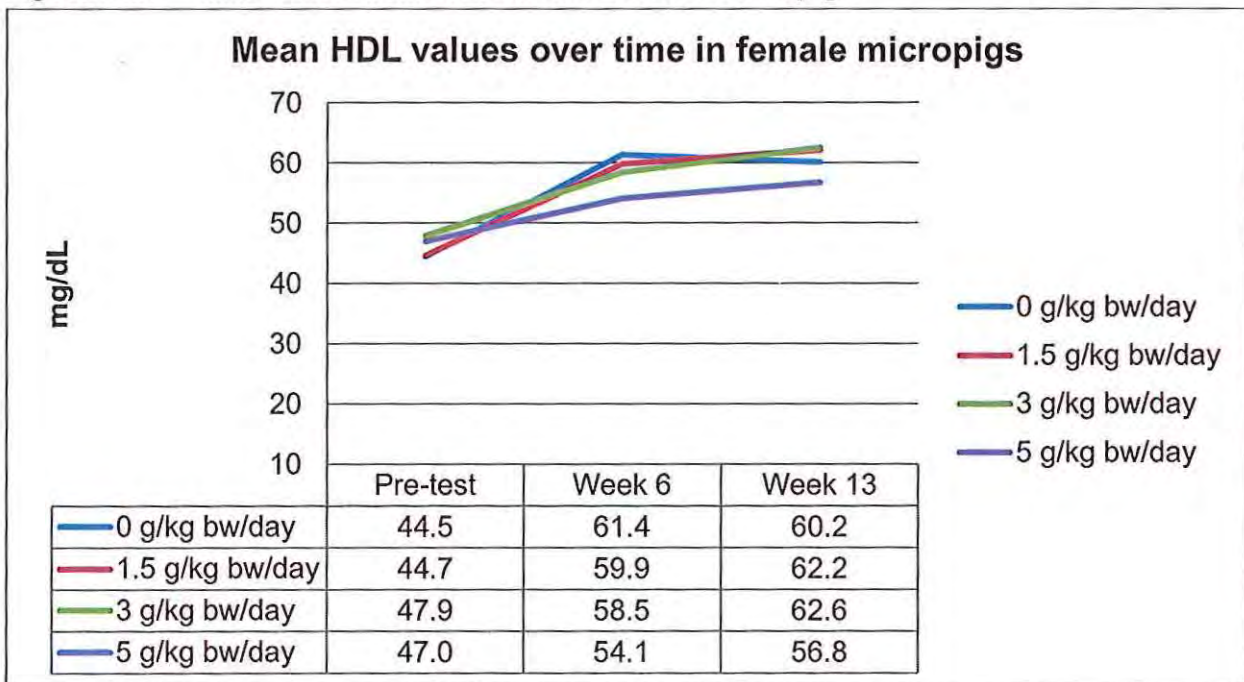


Figure 7-5b Mean HDL values over time in female micropigs

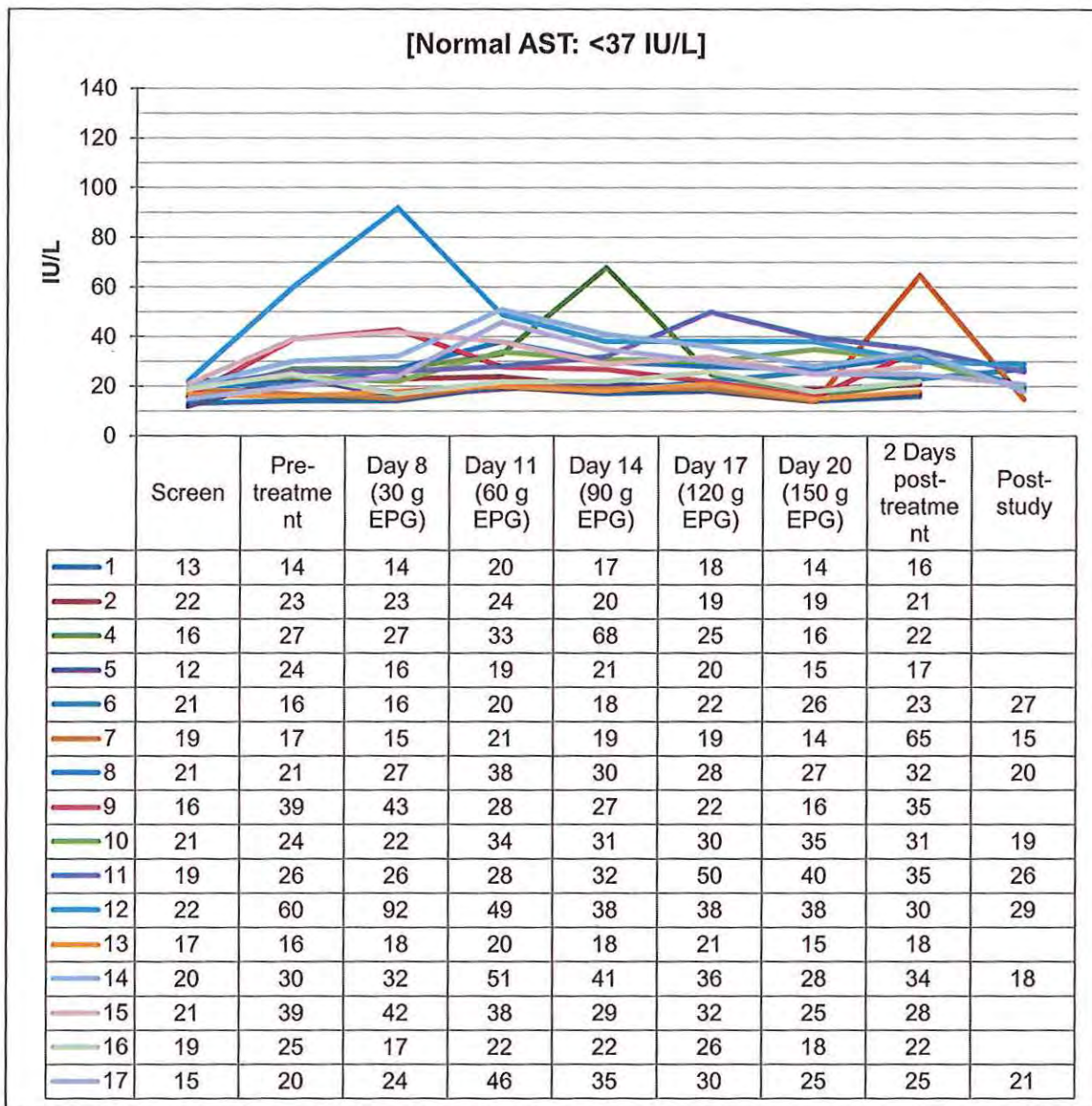


\* Values from 90-Day dietary toxicity study with esterified propoxylated glycerol (EPG) in Micropigs (Wedig and Bechtel, 2014)

## 7.5 Effect on Serum Transaminase Levels

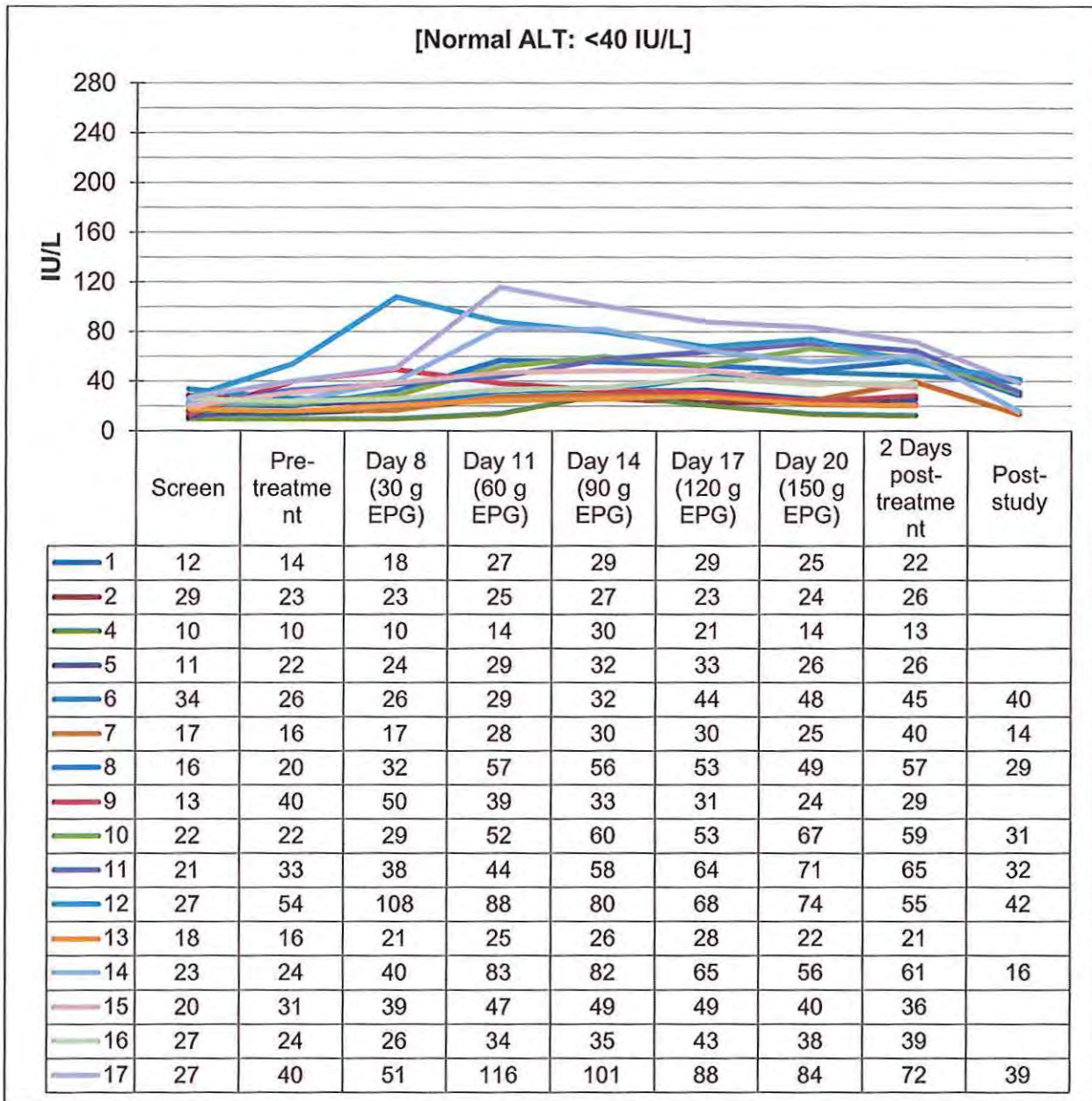
Results of multiple human range-finding tolerance studies showed that serum transaminase (aspartate aminotransferase and/or alanine aminotransaminase) levels exceeded the normal range in some subjects receiving EPG in amounts between 60 and 150 g/day (Figures 7-6a through 7-6d). The occasional moderate increase in measured serum transaminase values often occurred in a transient manner, rising briefly and then returning to normal ranges. This response was not observed in more extended clinical studies designed to also measure vitamin and nutrient status. Likewise, no effect on serum transaminase activity was reported in any of the animal studies including lifetime studies in rats and mice, nor was there reported any evidence of liver damage or related pathology in these investigations. Since the excursion of transaminase activity observed in some preliminary clinical investigations at doses exceeding 60 g/day, was not observed at lower doses or in extended preclinical safety investigations, it is not considered relevant to the much lower intake of EPG expected from the current intended uses.

**Figure 7-6a AST levels over time in each of 16 subjects receiving increasing EPG concentrations**



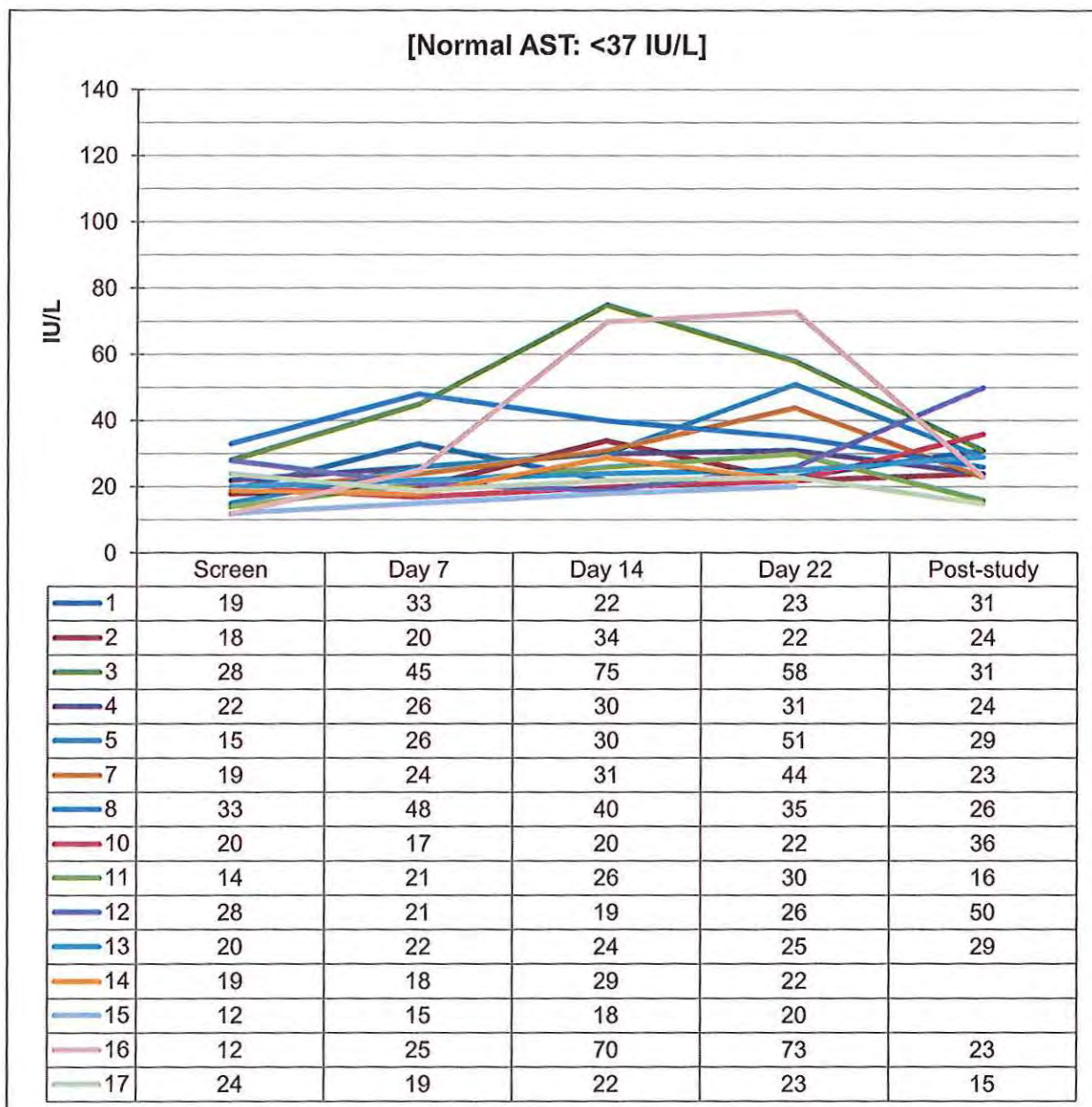
\* Values from ICR Project No. 0047047 (Section 6.1.1)

**Figure 7-6b ALT levels over time in each of 16 subjects receiving increasing EPG concentrations**



\* Values from ICR Project No. 0047047 (Section 6.1.1)

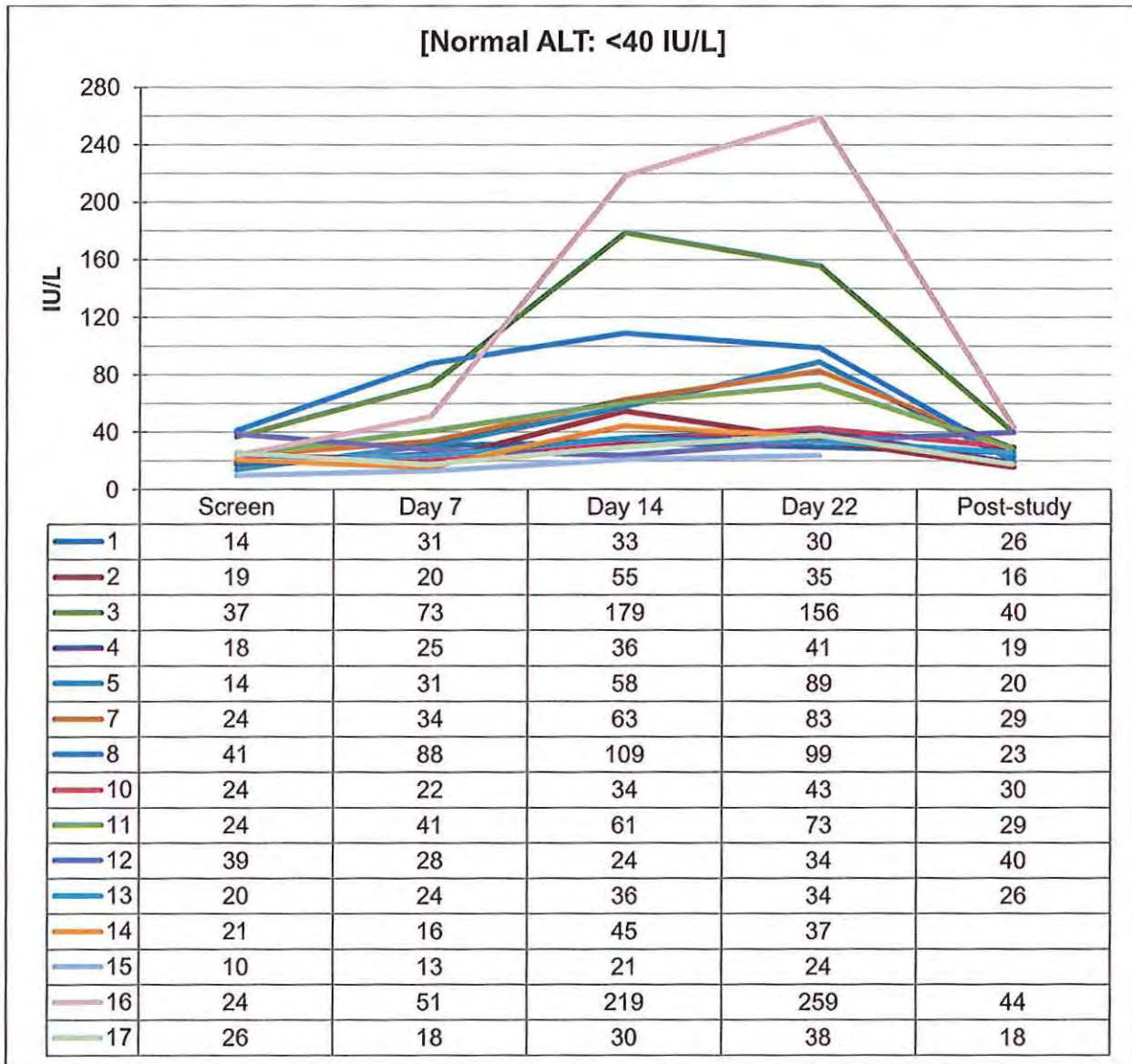
Figure 7-6c AST levels over time in each of 15 subjects receiving 120 g EPG/day



\* Values from ICR Project No. 004251 (Section 6.1.2)



Figure 7-6d ALT levels over time in each of 15 subjects receiving 120 g EPG/day



\* Values from ICR Project No. 004251 (Section 6.1.2)

## 8.0 SUMMARY AND CONCLUSIONS

This document demonstrates that use of Choco Finesse's H-EPG-05 as a fat replacer levels up to 38% (w/w expressed on a fat basis), in spreadable and baked goods such as, baked goods and baking mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, and soft candy, is GRAS.

The following constitute the basis for the GRAS determination:

- The structure of EPGs, being comprised of components of edible fats and oils interrupted by simple propylene glycol units, provides a basis for a strong presumption of safety. This presumption has been confirmed by the results of an extensive array of preclinical investigations, including lifetime feeding studies and those examining reproduction and developmental endpoint at doses up to 5 g EPG/kg body weight/day. No indication of toxicity was observed in any study.
- Low digestibility and caloric release was confirmed, as was lack of absorption of EPG following oral administration; EPG was not found in any tissue after up to a lifetime of feeding in rats.
- OSI test results demonstrated that EPG samples exhibit superior resistance to oxidation compared to shortenings commonly used in cooking and baking.
- Human and preclinical investigation of potential for nontoxic effects attributable to the physical state, consumed mass and solubility properties of EPG indicated low potential for significant or biologically meaningful effects at intake amounts anticipated for consumers. Gastrointestinal tolerance and tidiness were found to be augmented through selection of versions that are solid at human body temperature. Similarly, the potential for these untoward effects was minimized through selection of initial food applications that result in moderate consumer intake. Finally, studies in both experimental animals and humans demonstrated that: (1) the potential for interaction with lipid-soluble nutrients and other substances present in the gastrointestinal lumen is minimized through version selection (*i.e.*, a solid form of EPG) and moderation of consumption; and (2) there is low potential for biologically-meaningful effects at the maximum anticipated consumer intake. This is consistent with the moderate organic nature and solubility

properties of EPG (log Kow of approximately 3.2-3.4), and in strong contrast to the interaction of fat mimetics, such as olestra which have a log Kow<sup>16</sup> in excess of 40.

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<sup>16</sup> *n*-Octanol/water partition coefficient ( $K_{ow}$ ) is defined as the ratio of the molar concentrations of a chemical in *n*-octanol and water, in dilute solution. The coefficient  $K_{ow}$  is a constant for a given chemical at a given temperature (40 CFR 799.6756).

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## **APPENDIX 1**

**Estimated Daily Intake of Esterified Propoxylated Glycerols in Spreadable and Baked Food Products by the United States Population from Proposed Food-Uses**

## **Estimated Daily Intake of Esterified Propoxylated Glycerols in Spreadable and Baked Food Products by the United States Population from Proposed Food-Uses**

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# Estimated Daily Intake of Esterified Propoxylated Glycerols in Spreadable and Baked Food Products by the United States Population from Proposed Food-Uses

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# **Estimated Daily Intake of Esterified Propoxylated Glycerols in Spreadable and Baked Goods by the United States Population from Proposed Food-Uses**

## **1.0 INTRODUCTION**

Esterified propoxylated glycerols (EPG hereafter) are proposed for use in the United States (U.S.) in spreadable and baked goods, specifically baked goods and baking mixes, frozen dairy desserts and mixes, grain products and pasta, gravies and sauces, nuts and nut products, soft candy. EPG is intended for use to replace a portion of the fat content of foods and therefore, the percentage of proposed use level was adjusted for the fat content (%) of each intended food-use.

Estimates for the intake of EPG were based on the proposed food-uses and use-levels for EPG in conjunction with food consumption data included in the U.S. National Center for Health Statistics' (NCHS) National Health and Nutrition Examination Surveys (NHANES) 2011-2012 (CDC, 2014; USDA, 2014). Calculations for the mean and 90<sup>th</sup> percentile all-person and all-user intakes were performed for each of the individual proposed food-uses of EPG and the percentage of consumers were determined. Similar calculations were used to estimate the total intake of EPG resulting from all proposed food-uses of EPG combined. In both cases, the per person and per kilogram body weight intakes were reported for the following population groups:

- Infants and young children, 0 to 3 years;
- Children, ages 4 to 11;
- Female teenagers, ages 12 to 19;
- Male teenagers, ages 12 to 19;
- Female adults, ages 20 and up;
- Male adults, ages 20 and up; and
- Total population (all age and gender groups combined).

## **2.0 FOOD CONSUMPTION SURVEY DATA**

### **2.1 Survey Description**

NHANES for the years 2011-2012 are available for public use. NHANES are conducted as continuous, annual surveys, and are released in 2-year cycles. Each year about 7,000 people from 15 different locations across the U.S. are interviewed, and approximately 5,000 complete the health examination component of the survey. Any combination of consecutive years of data collection is recognized and used as a nationally representative sample of the U.S. population.

It is well-established that the length of a dietary survey affects the estimated consumption of individual users and that short-term surveys, such as a 1-day dietary survey, may overestimate consumption compared to surveys conducted over longer time periods (Anderson, 1988). Because two 24-hour dietary recalls administered on 2 non-consecutive days are available from the NHANES 2011-2012 survey, these data were used to generate estimates for the current intake analysis.

NHANES 2011-2012 survey data were collected from individuals and households *via* 24-hour dietary recalls administered on 2 non-consecutive days (Day 1 and Day 2) throughout all 4 seasons of the year. Day 1 data were collected in-person, and Day 2 data were collected by telephone in the following 3 to 10 days, on different days of the week, to achieve the desired degree of statistical independence. The data were collected by first selecting Primary Sampling Units (PSUs), which were counties throughout the U.S., of which 15 PSUs are visited per year. Small counties were combined to attain a minimum population size. These PSUs were segmented and households were chosen within each segment. One or more participants within a household were interviewed. For NHANES 2011-2012, 13,431 individuals were selected for the sample, 9,756 were interviewed (72.6%) and 9,338 were sampled (69.5%).

In addition to collecting information on the types and quantities of foods being consumed, NHANES 2011-2012 collected socio-economic, physiological and demographic information from individual participants in the survey, such as sex, age, height and weight, and other variables useful in characterizing consumption. The inclusion of this information allows for further assessment of food intake based on consumption by specific population groups of interest within the total population. The sample design for NHANES 2011-2012 includes an oversample of Asian Americans, however sample weights were incorporated to allow estimates from these subgroups to be combined to obtain national estimates that reflect the relative proportions of these groups in the population as a whole (CDC, 2014; USDA, 2014).

## 2.2 Statistical Methods

For the intake assessment, consumption data from individual dietary records, detailing food items ingested by each survey participant, were collated by computer and used to generate estimates for the intake of EPG by the U.S. population<sup>17</sup>. Estimates for the daily intake of EPG represent projected 2-day averages for each individual from Day 1 and Day 2 of NHANES 2011-2012 data; these average amounts comprised the distribution from which mean and percentile intake estimates were generated. Mean and percentile estimates were generated

---

<sup>17</sup> Statistical analysis and data management were conducted in DaDiet Software (Dazult Ltd., 2014). DaDiet Software is a web-based software tool that allows accurate estimate of exposure to nutrients and to EPGs added to foods, including contaminants, food additives and novel ingredients. The main input components are concentration (use level) data and food consumption data. Data sets are combined in the software to provide accurate and efficient exposure assessments.

incorporating survey weights in order to provide representative intakes for the entire U.S. population. All-person intake refers to the estimated intake of EPG averaged over all individuals surveyed, regardless of whether they potentially consumed food products containing EPG, and therefore includes individuals with “zero” intakes (*i.e.* those who reported no intake of food products containing EPG during the 2 survey days). All-user intake refers to the estimated intake of EPG by those individuals who reported consuming food products containing EPG, hence the “all-user” designation. Individuals were considered “users” if they consumed 1 or more food products containing EPG on either Day 1 or Day 2 of the survey.

Mean and 90<sup>th</sup> percentile intake estimates based on sample sizes of less than 30 and 80, respectively, may not be considered statistically reliable due to the limited sampling size (LSRO, 1995). As such, the reliability of estimates for the intake of EPG based on the consumption of these foods may be questionable for certain individual population groups. These values were not considered when assessing the relative contribution of specific food uses to total EPG consumption and are marked with an asterisk in Appendices A and B.

### **3.0 FOOD USAGE DATA**

The individual proposed food-uses and use-levels for EPG employed in the current intake analysis are summarized in Table 3-1. Food codes representative of each proposed food-use were chosen from the NHANES 2011-2012 (CDC, 2014; USDA, 2014). Food codes were grouped in food-use categories according to Title 21, Section §170.3 of the Code of Federal Regulations (CFR, 2014a). Product-specific adjustment factors were developed based on data provided in the standard recipe file for the Continuing Survey of Food Intakes by Individuals (CSFII) 1994-1996, 1998 survey (USDA, 2000). All food codes included in the current intake assessment are listed in Appendix C.

**Table 3-1 Summary of the Individual Proposed Uses and Use Levels for EPG in the United States (2011-2012 NHANES Data)**

Food Category	Proposed Food-Uses	Percent (%) Fat in Product <sup>a</sup>	Percent (%) EPG Inclusion Expressed on Fat Basis	Percent (%) EPG Inclusion in Final Product <sup>b</sup>
Baked goods and baking mixes	Biscuits	14 to 18	4-12	2.16
	Breads, specialty, flavored	7 to 11	2-8	0.88
	Brownies	22 to 30	6-17	5.10
	Cakes	15 to 20	4-14	2.80
	Cookies	22 to 30	7-21	6.30
	Crepes	8 to 28	2-20	5.60
	Desserts excluding cakes, cookies, brownies	8 to 28	2-20	5.60
	Doughnuts	20 to 23	5-12	2.76
	Muffins	9 to 19	2-13	2.47
	Pastry	21	6-15	3.15
	Pastry crusts and pies	23 to 36	7-25	9.00
Frozen dairy desserts and mixes	Ice cream	15 to 21	4-12	2.52
Grain products and pastas	Granola and other snack bars (e.g., pumpkin-based) <sup>c</sup>	14 to 24	4-17	4.08
Gravies and sauces	Pasta sauces (cream and tomato based)	11	3-8	0.88
Nut and nut products	Nut butters and nut spreads	32 to 55	10-38	20.90
Soft candy	Candy bars	14 to 24	4-17	4.08

<sup>a</sup> Values listed were provided by the client. These were compared to the mean values per category available in the 2011-2012 NHANES data and noted to be higher; therefore, the intake assessment conducted using the percentage of fat provided by the client presented herein is considered to be a conservative estimate.

<sup>b</sup> Calculated by (Maximum percent fat in product) \* (Maximum percent EPG inclusion expressed on a fat basis)

<sup>c</sup> No pumpkin-based bars were identified in the NHANES dataset, therefore this food use has not been included in the present assessment.

## 4.0 FOOD SURVEY RESULTS

Estimates for the total daily intakes of EPG from proposed food-uses are provided in Tables 4.1-1 and 4.1-2. Estimates for the daily intake of EPG from individual proposed food-uses in the U.S. are summarized in Tables A-1 to A-7 and B-1 to B-7 of Appendices A and B, respectively. Tables A-1 to A-7 provide estimates for the daily intake of EPG per person (g/day), whereas Tables B-1 to B-7 provide estimates for the daily intake of EPG on a per kilogram body weight basis (mg/kg body weight/day).

## 4.1 Estimated Daily Intake of EPG from All Proposed Food-Uses in the United States

Table 4.1-1 summarizes the estimated total intake of EPG (g/person/day) from all proposed food-uses in the U.S. population group. Table 4.1-2 presents this data on a per kilogram body weight basis (mg/kg body weight/day). The percentage of users was high among all age groups evaluated in the current intake assessment; greater than 70.3% of the population groups consisted of users of those food products in which EPG is currently proposed for use (Table 4.1-1). Children had the greatest percentage of users at 91.7%. Large user percentages within a population group typically lead to similar results for the all-person and all-user consumption estimates. Consequently, only the all-user intake results will be discussed in detail.

Among the total population, the mean and 90<sup>th</sup> percentile all-user intakes of EPG were determined to be 3.6 and 8.0 g/person/day, respectively. Of the individual population groups, male adults were determined to have the greatest mean and 90<sup>th</sup> percentile all-user intakes of EPG on an absolute basis, at 4.1 and 9.2 g/person/day, respectively, while infants had the lowest mean and 90<sup>th</sup> percentile all-user intakes of 2.3 and 5.2 g/person/day, respectively (Table 4.1-1).

**Table 4.1-1 Summary of the Estimated Daily Intake of EPG from Proposed Food-Uses in the United States by Population Group (2011-2012 NHANES Data)**

Population Group	Age Group (Years)	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	% Users	n	Mean	90 <sup>th</sup> Percentile
Infants and Young Children	0 to 3	1.6	4.2	70.3	547	2.3	5.2
Children	4 to 11	3.2	7.1	91.7	1,198	3.5	7.4
Female Teenagers	12 to 19	2.2	5.5	77.6	429	2.8	5.7
Male Teenagers	12 to 19	3.0	7.7	78.3	419	3.9	9.0
Female Adults	20 and up	2.7	6.8	82.6	1,761	3.3	7.3
Male Adults	20 and up	3.4	8.7	82.5	1,645	4.1	9.2
Total Population	All Ages	2.9	7.3	82.4	5,999	3.6	8.0

On a body weight basis, infants and young children were identified as having the highest mean and 90<sup>th</sup> percentile all-user intakes of any population group, of 168 and 358 mg/kg body weight/day, respectively. Female adults had the lowest mean and 90<sup>th</sup> percentile all-user intakes of 46 and 104 mg/kg body weight/day, respectively (Table 4.1-2).



**Table 4.1-2 Summary of the Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Proposed Food-Uses in the United States by Population Group (2011-2012 NHANES Data)**

Population Group	Age Group (Years)	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
Infants and Young Children	0 to 3	118	304	70.4	546	168	358
Children	4 to 11	115	277	91.7	1,198	125	280
Female Teenagers	12 to 19	39	102	77.2	419	50	120
Male Teenagers	12 to 19	49	130	78.2	416	63	153
Female Adults	20 and up	38	93	82.6	1,746	46	104
Male Adults	20 and up	40	103	82.5	1,630	49	107
Total Population	All Ages	52	128	82.4	5,955	63	146

## 4.2 Estimated Daily Intake of EPG from Individual Proposed Food-Uses in the United States

Estimates for the mean and 90<sup>th</sup> percentile daily intakes of EPG from each individual food category are summarized in Tables A-1 to A-7 and B-1 to B-7 on a g/day and mg/kg body weight/day basis, respectively. The total U.S. population was identified as being significant consumers of cookies (26.4 to 46.8% users), ice cream (15.4 to 33.3% users) and pasta sauces (19.9 to 27.8% users).

In terms of contribution to total mean intake of EPG, cookies (contributed 19.5 to 31.2% to total mean intakes) and nut butters (contributed 16.5 to 28.3% to total mean intakes) were the 2 main sources of intake across all population groups on both an absolute and on a mg/kg body weight basis. Pastries individually contributed  $\leq 1.0\%$  to total mean EPG intakes across all population groups (see Tables A-1 to A-1 and/or B-1 to B-7 for further details).

## 5.0 CONCLUSIONS

Consumption data and information pertaining to the individual proposed food-uses of EPG were used to estimate the all-person and all-user intakes of EPG for specific demographic groups and for the total U.S. population. There were a number of assumptions included in the assessment which mean that the resulting exposure estimates may be considered the “worst case” values. For example, it has been assumed in both exposure assessments that all food products within a food category contain EPG at the maximum specified level of use. In reality, the levels added to specific foods will vary. In addition, it is well-established that the length of a dietary survey affects the estimated consumption of individual users. Short-term surveys, such as the typical 2-

or 3-day dietary surveys, may overestimate the consumption of food products that are consumed relatively infrequently (Anderson, 1988).

In summary, on an all-user basis, the resulting mean and 90<sup>th</sup> percentile intakes of EPG by the total U.S. population from all proposed food-uses in the U.S., were estimated to be 3.6 g/person/day (63 mg/kg body weight/day) and 8.0 g/person/day (146 mg/kg body weight/day), respectively. Among the individual population groups, the highest mean and 90<sup>th</sup> percentile intakes of EPG were determined to be 4.1 g/person/day (49 mg/kg body weight/day) and 9.2 g/person/day (107 mg/kg body weight/day), respectively, as identified among male adults. When expressed on a body weight basis, infants and young children were identified as having the highest mean and 90<sup>th</sup> percentile all-user intakes of any population group, of 168 and 358 mg/kg body weight/day, respectively.

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**Appendix A**  
**Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Different**  
**Population Groups Within the United States (2011-2012 NHANES DATA)**

**Table A-1 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Infants and Young Children Aged 0 to 3 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	1.6	4.2	70.3	547	2.3	5.2
<u>Baked Goods and Baking Mixes</u>							
Biscuits	1.6	0.03	na	4.7	37	0.56	0.84*
Breads, specialty, flavored	1.0	0.02	na	8.6	64	0.19	0.37*
Brownies	<0.1	<0.01*	na	0.5	6	0.86*	0.77*
Cakes	4.5	0.07	na	6.6	64	1.11	2.92*
Cookies	31.2	0.51	1.62	36.3	296	1.40	2.71
Crepes	0	na	na	0	0	na	na
Desserts excluding cakes, cookies, brownies	<0.1	<0.01*	na	<0.1	1	0.19*	0.17*
Doughnuts	1.8	0.03	na	5.6	32	0.53	1.05*
Muffins	1.9	0.03*	na	2.3	20	1.39*	1.61*
Pastry	<0.1	<0.01*	na	0.3	3	0.61*	0.64*
Pastry crusts and pies	3.0	0.05*	na	2.1	9	2.27*	3.31*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	9.0	0.15	0.71	15.4	114	0.96	1.71
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	6.4	0.10	0.38*	11	48	0.94	1.68*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.5	0.06	0.21	21.7	154	0.27	0.63
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	28.3	0.46	1.67	20.5	119	2.25	5.02
<u>Soft Candy</u>							
Candy bars	7.2	0.12	0.13*	11.5	73	1.02	3.22*

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table A-2 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Children Aged 4 to 11 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<b>All</b>	<b>100</b>	3.2	7.1	91.7	1,198	3.5	7.4
<b>Baked Goods and Baking Mixes</b>							
Biscuits	2.4	0.08	na	8.8	98	0.87	1.52
Breads, specialty, flavored	1.5	0.05	0.13	12.1	154	0.40	0.80
Brownies	2.3	0.07	na	6	69	1.23	2.30*
Cakes	7.5	0.24	0.87	17	214	1.39	3.29
Cookies	27.7	0.88	2.62	46.8	560	1.88	3.78
Crepes	<0.1	<0.01*	na	<0.1	2	3.87*	4.14*
Desserts excluding cakes, cookies, brownies	0.4	0.01*	na	0.5	8	2.81*	3.22*
Doughnuts	2.2	0.07	na	9.1	106	0.76	1.28
Muffins	3.7	0.12	na	7.2	72	1.62	2.78*
Pastry	<0.1	<0.01*	na	0.4	6	0.65*	0.65*
Pastry crusts and pies	3.5	0.11	na	2.3	32	4.79	7.63*
<b>Frozen Dairy Desserts and Mixes</b>							
Ice Cream	16.8	0.53	1.87	33.3	406	1.60	3.01
<b>Grain Products and Pastas</b>							
Granola and other snack bars (e.g., pumpkin-based)	3.7	0.12	0.71	13.7	141	0.86	1.39
<b>Gravies and Sauces</b>							
Pasta sauces (cream and tomato based)	3.3	0.10	0.33	27.8	364	0.37	0.85
<b>Nut and Nut Products</b>							
Nut butters and nut spreads	21.5	0.68	2.23	23.3	275	2.91	6.69
<b>Soft Candy</b>							
Candy bars	3.4	0.11	0.38	17.4	229	0.62	1.18

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table A-3 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Female Teenagers Aged 12 to 19 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	2.2	5.5	77.6	429	2.8	5.7
<u>Baked Goods and Baking Mixes</u>							
Biscuits	3.5	0.07	na	9	41	0.83	1.38*
Breads, specialty, flavored	1.7	0.04	0.14*	10.9	44	0.34	0.64*
Brownies	2.1	0.05*	na	3	20	1.51*	3.00*
Cakes	9.8	0.21	<0.01*	10.1	68	2.11	5.71*
Cookies	19.8	0.43	1.89	26.4	167	1.63	2.84
Crepes	<0.1	<0.01*	na	<0.1	2	3.24*	4.26*
Desserts excluding cakes, cookies, brownies	0.7	0.02*	na	0.9	3	1.75*	1.85*
Doughnuts	3.0	0.07*	na	6.4	25	1.01*	1.49*
Muffins	2.6	0.06*	na	4.5	19	1.25*	1.70*
Pastry	<0.1	<0.01*	na	0.1	2	1.42*	1.59*
Pastry crusts and pies	3.4	0.07*	na	1.7	12	4.36*	6.51*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	21.1	0.46	1.63	23.5	145	1.94	3.73
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	5.5	0.12	0.43*	11.7	64	1.01	1.75*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	4.1	0.09	0.20	20.2	110	0.43	1.30
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	16.5	0.36	1.67*	12.6	50	2.82	5.02*
<u>Soft Candy</u>							
Candy bars	6.1	0.13	0.42	16.8	89	0.79	1.61

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table A-4 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Male Teenagers Aged 12 to 19 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	3.0	7.7	78.3	419	309	9.0
<u>Baked Goods and Baking Mixes</u>							
Biscuits	3.3	0.10	na	9	44	1.12	3.11*
Breads, specialty, flavored	1.4	0.04	na	8.5	36	0.50	0.65*
Brownies	1.4	0.04*	na	3.4	23	1.21*	1.66*
Cakes	7.1	0.21	na	9.6	59	2.23	3.43*
Cookies	19.6	0.59	1.89	29.4	190	2.02	4.51
Crepes	0	0	0	0	0	0	0
Desserts excluding cakes, cookies, brownies	0.7	0.02*	na	0.6	3	3.62*	3.73*
Doughnuts	3.3	0.10	na	8.2	40	1.23	2.65*
Muffins	1.6	0.05*	na	3	16	1.55*	2.22*
Pastry	<0.1	<0.01*	na	0.1	2	0.89*	1.14*
Pastry crusts and pies	12.2	0.37*	na	4.4	15	8.35*	9.86*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	16.9	0.51	1.87	21.1	109	2.43	4.54
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	4.0	0.12	0.49*	11.5	48	1.06	1.44*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.9	0.12	0.35	19.9	105	0.59	1.44
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	19.6	0.59	3.34*	16.8	54	3.54	6.69*
<u>Soft Candy</u>							
Candy bars	4.9	0.15	0.47*	14.3	66	1.04	2.37*

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Table A-5 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Female Adults Aged 20 Years and Over Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	2.7	6.8	82.6	1,761	3.3	7.3
<u>Baked Goods and Baking Mixes</u>							
Biscuits	2.0	0.05	na	6.4	165	0.86	1.49
Breads, specialty, flavored	1.3	0.04	na	9.7	223	0.36	0.65
Brownies	1.7	0.05	na	4.1	75	1.13	1.66*
Cakes	10.2	0.28	1.10	17.3	382	1.61	2.99
Cookies	19.5	0.53	1.89	31.6	672	1.69	3.28
Crepes	0.3	0.01*	na	0.2	7	3.24*	4.37*
Desserts excluding cakes, cookies, brownies	2.6	0.07	na	2	33	3.63	6.85*
Doughnuts	1.6	0.04	na	4.6	100	0.97	2.00
Muffins	2.0	0.05	na	3.8	71	1.43	3.21*
Pastry	0.8	0.02*	na	0.8	24	2.86*	5.09*
Pastry crusts and pies	14.5	0.40	na	7	117	5.62	8.35
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	13.8	0.38	1.51	21.6	468	1.75	3.38
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.8	0.10	0.43	10.3	184	1.01	1.71
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.1	0.08	0.28	21.1	421	0.40	0.91
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	17.3	0.47	1.67	15.9	266	2.97	6.69
<u>Soft Candy</u>							
Candy bars	5.5	0.15	0.55	18.5	345	0.80	1.37

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table A-6 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by Male Adults Aged 20 Years and Over Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<b>All</b>	<b>100</b>	3.4	8.7	82.5	1,645	4.1	9.2
<u>Baked Goods and Baking Mixes</u>							
Biscuits	3.1	0.11	0.32	10.2	206	1.03	1.80
Breads, specialty, flavored	1.0	0.03	na	6.9	145	0.49	0.98
Brownies	1.3	0.04	na	3.2	51	1.35	2.30*
Cakes	8.4	0.29	1.10	15	287	1.90	3.74
Cookies	21.4	0.73	2.77	32.2	658	2.27	4.54
Crepes	0.2	0.01*	na	0.1	3	5.61*	4.64*
Desserts excluding cakes, cookies, brownies	1.7	0.06*	na	1.8	27	3.19*	4.72*
Doughnuts	2.1	0.07	na	6.2	114	1.15	1.96
Muffins	2.0	0.07	na	3.6	59	1.88	3.21*
Pastry	0.6	0.02*	na	1.4	26	1.50*	2.05*
Pastry crusts and pies	13.5	0.46	na	6	111	7.71	13.74
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	14.4	0.49	1.70	22.5	448	2.19	4.54
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.5	0.12	na	9.8	140	1.23	2.45
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.7	0.13	0.38	22	427	0.57	1.44
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	18.3	0.63	1.67	13.9	241	4.50	10.03
<u>Soft Candy</u>							
Candy bars	4.9	0.17	0.60	17.7	319	0.94	1.74

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table A-7 Estimated Daily Intake of EPG from Individual Proposed Food-Uses by the Total United States Population (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (g/day)		All-Users Consumption (g/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	2.9	7.3	82.4	5,999	3.6	8.0
<u>Baked Goods and Baking Mixes</u>							
Biscuits	2.6	0.08	na	8.2	591	0.94	1.56
Breads, specialty, flavored	1.2	0.04	na	8.9	666	0.40	0.71
Brownies	1.6	0.05	na	3.7	244	1.23	1.81
Cakes	8.8	0.26	1.05	15.1	1,074	1.71	3.43
Cookies	21.6	0.64	2.21	33.3	2,543	1.91	3.78
Crepes	<0.1	<0.01*	na	0.1	14	3.96*	4.53*
Desserts excluding cakes, cookies, brownies	1.7	0.05	na	1.5	75	3.35	6.48*
Doughnuts	2.0	0.06	na	6	417	1.00	1.96
Muffins	2.2	0.06	na	4	257	1.60	3.21
Pastry	0.6	0.02	na	0.9	63	1.91	2.68*
Pastry crusts and pies	11.9	0.35	na	5.5	296	6.42	12.02
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	14.7	0.43	1.63	22.9	1,690	1.90	3.96
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.8	0.11	0.43	10.7	625	1.06	1.75
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.5	0.10	0.31	22	1,581	0.46	1.14
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	18.6	0.55	1.67	16.1	1,005	3.40	6.69
<u>Soft Candy</u>							
Candy bars	5.0	0.15	0.53	17.4	1,121	0.85	1.71

na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Appendix B**  
**Estimated Daily Per Kilogram Body Weight Intake of EPG from Individual**  
**Proposed Food-Uses by Different Population Groups Within the United States**  
**(2011-2012 NHANES DATA)**

**Table B-1 Estimated Daily Per Kilogram Body Weight Intake of EPG from Individual Proposed Food-Uses by Infants and Young Children Aged 0 to 3 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	n	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	118	304	70.4	546	168	358
<u>Baked Goods and Baking Mixes</u>							
Biscuits	1.5	2	na	4.7	37	38	50*
Breads, specialty, flavored	1.1	1	na	8.6	64	15	29*
Brownies	0.3	<1*	na	0.5	6	56*	53*
Cakes	4.3	5	na	6.6	64	77	190*
Cookies	31.5	37	118	36.3	295	102	187
Crepes	0	0	0	0	0	0	0
Desserts excluding cakes, cookies, brownies	<0.1	<1*	na	<0.1	1	17*	15*
Doughnuts	1.7	2	na	5.6	32	35	85*
Muffins	1.8	2*	na	2.3	20	92*	106*
Pastry	0.1	<1*	na	0.3	3	34*	35*
Pastry crusts and pies	2.8	3*	na	2.1	9	153*	196*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	8.7	10	53	15.4	114	67	120
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	6.4	8	26*	11	48	69	123*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.7	4	15	21.7	154	20	50
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	29.3	35	142	20.5	119	169	353
<u>Soft Candy</u>							
Candy bars	6.9	8	9*	11.5	73	71	190*

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table B-2 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by Children Aged 4 to 11 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<b>All</b>	<b>100</b>	115	277	91.7	1,198	125	280
<u>Baked Goods and Baking Mixes</u>							
Biscuits	2.4	3	na	8.8	98	31	70
Breads, specialty, flavored	1.5	2	5	12.1	154	15	28
Brownies	2.3	3	na	6	69	45	95*
Cakes	7.2	8	30	17	214	48	117
Cookies	27.4	32	97	46.8	560	67	128
Crepes	<0.1	<1*	na	<0.1	2	149*	168*
Desserts excluding cakes, cookies, brownies	0.4	1*	na	0.5	8	106*	161*
Doughnuts	2.2	3	na	9.1	106	28	48
Muffins	3.7	4	na	7.2	72	59	102*
Pastry	0.1	<1*	na	0.4	6	30*	30*
Pastry crusts and pies	3.1	4	na	2.3	32	156	295*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	15.9	18	62	33.3	406	55	114
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.8	4	20	13.7	141	32	54
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.3	4	14	27.8	364	14	36
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	23.5	27	95	23.3	275	116	221
<u>Soft Candy</u>							
Candy bars	3.1	4	13	17.4	229	20	42

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table B-3 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by Female Teenagers Aged 12 to 19 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<i>All</i>	100	39	102	77.2	419	50	120
<u>Baked Goods and Baking Mixes</u>							
Biscuits	2.4	1	na	8.3	37	11	15*
Breads, specialty, flavored	1.5	1	2*	10.6	42	5	9*
Brownies	2.3	1*	na	3.1	20	28*	57*
Cakes	10.3	4	na	9.9	67	40	113*
Cookies	21.2	8	34	26.9	166	30	66
Crepes	0.1	<1*	na	<0.1	2	56*	77*
Desserts excluding cakes, cookies, brownies	0.7	<1*	na	0.9	3	30*	33*
Doughnuts	2.8	1*	na	6.6	24	17*	28*
Muffins	3.4	1*	na	4.6	19	28*	37*
Pastry	<0.1	<1*	na	0.1	2	19*	20*
Pastry crusts and pies	3.2	1*	na	1.6	11	79*	114*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	21.5	8	31	23.3	143	36	61
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	5.9	2	8*	11.9	64	19	35*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	4.1	2	4	20.3	107	8	24
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	15.5	6	28*	12.7	49	47	105*
<u>Soft Candy</u>							
Candy bars	5.2	2	8	16.9	86	12	25

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Table B-4 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by Male Teenagers Aged 12 to 19 Years Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<i>All</i>	100	49	130	78.2	416	63	153
<u>Baked Goods and Baking Mixes</u>							
Biscuits	3.4	2	na	9	44	19	42*
Breads, specialty, flavored	1.3	1	na	8.3	34	8	12*
Brownies	1.1	1*	na	3.4	23	16*	20*
Cakes	6.5	3	na	9.6	58	33	54*
Cookies	20.0	10	33	29.5	189	33	74
Crepes	0	Na	na	0	0	na	na
Desserts excluding cakes, cookies, brownies	0.6	<1*	na	0.6	3	49*	50*
Doughnuts	3.4	2	na	8.2	40	20	37*
Muffins	1.8	1*	na	3	16	28*	45*
Pastry	<0.1	<1*	na	0.1	2	13*	14*
Pastry crusts and pies	13.0	6*	na	4.4	15	144*	202*
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	16.8	8	27	21.1	108	39	94
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	4.4	2	9*	11.5	48	19	37*
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.5	2	6	20	104	9	19
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	19.4	9	44*	16.8	54	57	105*
<u>Soft Candy</u>							
Candy bars	4.7	2	6*	14.3	66	16	35*

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Table B-5 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by Female Adults Aged 20 Years and Over Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<b>All</b>	<b>100</b>	38	93	82.6	1,746	46	104
<u>Baked Goods and Baking Mixes</u>							
Biscuits	1.9	1	na	6.3	163	11	19
Breads, specialty, flavored	1.2	<1	na	9.6	220	5	8
Brownies	2.0	1	na	4.1	75	18	26*
Cakes	10.2	4	16	17.2	376	23	47
Cookies	19.8	8	27	31.5	666	24	51
Crepes	0.3	<1*	na	0.2	7	53*	70*
Desserts excluding cakes, cookies, brownies	2.4	1	na	2	32	47	102*
Doughnuts	1.7	1	na	4.6	100	14	25
Muffins	2.1	1	na	3.8	71	21	44*
Pastry	0.9	<1*	na	0.8	24	42*	84*
Pastry crusts and pies	14.5	6	na	7.1	116	78	140
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	13.1	5	19	21.7	465	23	46
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.7	1	4	10.4	183	14	25
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.2	1	4	21	418	6	15
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	17.5	7	24	15.8	262	43	83
<u>Soft Candy</u>							
Candy bars	5.5	2	7	18.4	343	11	21

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Table B-6 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by Male Adults Aged 20 Years and Over Within the United States (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	40	103	82.5	1,630	49	107
<u>Baked Goods and Baking Mixes</u>							
Biscuits	3.0	1	3	10.2	206	12	20
Breads, specialty, flavored	1.0	<1	na	7	145	6	11
Brownies	1.3	1	na	3.2	50	16	28*
Cakes	8.0	3	12	14.9	282	22	42
Cookies	21.6	9	33	32.3	651	27	55
Crepes	0.2	<1*	na	0.1	3	74*	66*
Desserts excluding cakes, cookies, brownies	1.6	1*	na	1.8	27	36*	59*
Doughnuts	2.0	1	na	6.2	114	13	23
Muffins	1.9	1	na	3.6	59	21	37*
Pastry	0.6	<1*	na	1.4	26	17*	23*
Pastry crusts and pies	13.2	5	na	6	111	88	157
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	14.6	6	21	22.6	445	26	60
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	3.5	1	na	9.8	138	14	27
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.8	2	4	22.1	427	7	17
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	18.8	8	23	14	239	54	112
<u>Soft Candy</u>							
Candy bars	4.9	2	7	17.7	317	11	22

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.

**Table B-7 Estimated Daily Per Kilogram Body Weight Intake of Esterified Propoxylated Glycerol from Individual Proposed Food-Uses by the Total United States Population (2011-2012 NHANES Data)**

Food-Use Category	% Contribution to Total Mean Intake	All-Person Consumption (mg/kg bw/day)		All-Users Consumption (mg/kg bw/day)			
		Mean	90 <sup>th</sup> Percentile	%	N	Mean	90 <sup>th</sup> Percentile
<i>All</i>	<i>100</i>	52	128	82.4	5,955	63	146
<u>Baked Goods and Baking Mixes</u>							
Biscuits	2.4	1	na	8.1	585	15	32
Breads, specialty, flavored	1.2	1	na	8.9	659	7	14
Brownies	1.6	1	na	3.7	243	23	44
Cakes	8.0	4	14	15	1,061	28	54
Cookies	23.5	12	39	33.3	2,527	37	78
Crepes	0.2	<1*	na	0.1	14	62*	73*
Desserts excluding cakes, cookies, brownies	1.3	1	na	1.5	74	44	105*
Doughnuts	2.0	1	na	6	416	18	36
Muffins	2.4	1	na	4	257	31	70
Pastry	0.5	<1	na	0.9	63	27	37*
Pastry crusts and pies	9.5	5	na	5.5	294	90	169
<u>Frozen Dairy Desserts and Mixes</u>							
Ice Cream	14.2	7	25	23	1,681	32	67
<u>Grain Products and Pastas</u>							
Granola and other snack bars (e.g., pumpkin-based)	4.1	2	6	10.7	622	20	41
<u>Gravies and Sauces</u>							
Pasta sauces (cream and tomato based)	3.5	2	5	22.1	1,574	8	20
<u>Nut and Nut Products</u>							
Nut butters and nut spreads	20.8	11	31	16.1	998	67	146
<u>Soft Candy</u>							
Candy bars	4.9	3	8	17.4	1,114	15	28

bw = body weight; na = not available

\* Indicates an intake estimate that may not be statistically reliable, as the sample size does not meet the minimum reporting requirements.



**Appendix C**  
**Representative NHANES Food Codes for Proposed Food-Uses of Esterified Propoxylated Glycerol in the United States (2011-2012 NHANES DATA)**

## Representative NHANES Food Codes for Proposed Food-Uses of Esterified Propoxylated Glycerol (EPG) in the United States

### Baked Goods and Baking Mixes

#### Biscuits

[EPG] = 0.02% (12% adjusted for fat portion of 18%)

52101000	Biscuit, baking powder or buttermilk type, NS as to made from mix, refrigerated dough, or home recipe
52101020	Biscuit dough, raw
52101030	Biscuit dough, fried
52101100	Biscuit, baking powder or buttermilk type, made from mix
52101150	Biscuit, baking powder or buttermilk type, made from refrigerated dough, lowfat
52102040	Biscuit, baking powder or buttermilk type, made from refrigerated dough
52103000	Biscuit, baking powder or buttermilk type, commercially baked
52104040	Biscuit, whole wheat
52104100	Biscuit, cheese
52104200	Biscuit, cinnamon-raisin

#### Mixed foods containing biscuits

*Adjusted for biscuits content of 68 to ~90%*

27520170	Bacon on biscuit
27520250	Ham on biscuit
27540145	Chicken fillet (breaded, fried) sandwich on biscuit
27540180	Chicken patty sandwich or biscuit
27560650	Sausage on biscuit
32202020	Egg, cheese, and ham on biscuit
27515080	Steak sandwich, plain, on biscuit

*Adjusted for biscuits content of 30 to 57%*

27515080	Steak sandwich, plain, on biscuit
32202060	Egg and sausage on biscuit
32202090	Egg and bacon on biscuit
32202110	Egg and ham on biscuit
32202130	Egg and steak on biscuit
32202070	Egg, cheese, and bacon on biscuit
32202200	Egg and cheese on biscuit
32202050	Egg, cheese, and sausage on biscuit
58128000	Biscuit with gravy

#### Breads, Specialty, Flavored

[EPG] = 0.01% (11% adjusted for fat portion of 11%)

51111010	Bread, cheese
51111040	Bread, cheese, toasted
51113010	Bread, cinnamon
51113100	Bread, cinnamon, toasted
51115010	Bread, cornmeal and molasses
51115020	Bread, cornmeal and molasses, toasted
51119010	Bread, egg, Challah

51119040	Bread, egg, Challah, toasted
51121010	Bread, garlic
51121110	Bread, onion
51121120	Bread, onion, toasted
51127010	Bread, potato
51127020	Bread, potato, toasted
51129010	Bread, raisin
51129020	Bread, raisin, toasted
51134000	Bread, sweet potato
51134010	Bread, sweet potato, toasted
51135000	Bread, vegetable
51135010	Bread, vegetable, toasted
51300210	Bread, whole wheat, with raisins
51300220	Bread, whole wheat, with raisins, toasted
51301120	Bread, wheat or cracked wheat, with raisins
51301130	Bread, wheat or cracked wheat, with raisins, toasted
51601210	Bread, multigrain, with raisins
51601220	Bread, multigrain, with raisins, toasted
52201000	Cornbread, prepared from mix
52401000	Bread, Boston Brown
52403000	Bread, nut
52404060	Bread, pumpkin
52405010	Bread, fruit
52407000	Bread, zucchini
52408000	Bread, Irish soda
55301000	French toast, plain
55301050	French toast sticks, plain

## Brownies

[EPG] = 0.05 (17% adjusted for fat portion of 30%)

53204000	Cookie, brownie, NS as to icing
53204010	Cookie, brownie, without icing
53204100	Cookie, brownie, with icing
53204840	Cookie, brownie, reduced fat, NS as to icing
53204860	Cookie, brownie, fat free, without icing
53205250	Cookie, butterscotch, brownie

## Cakes

[EPG] = 0.03 (14% adjusted for fat portion of 20%)

53100100	Cake or cupcake, NS as to type
53101100	Cake, angel food, without icing or filling
53101200	Cake, angel food, with icing or filling
53101250	Cake, angel food, with fruit and icing or filling
53102100	Cake or cupcake, applesauce, without icing or filling
53102200	Cake or cupcake, applesauce, with icing or filling
53102600	Cake or cupcake, banana, without icing or filling
53102700	Cake or cupcake, banana, with icing or filling
53102800	Cake or cupcake, black forest (chocolate-cherry)
53103000	Cake, Boston cream pie
53104100	Cake or cupcake, carrot, without icing or filling
53104260	Cake or cupcake, carrot, with icing or filling
53104300	Cake, carrot, diet
53104400	Cake or cupcake, coconut, with icing or filling

53104500	Cheesecake
53104550	Cheesecake with fruit
53104600	Cheesecake, chocolate
53105270	Cake or cupcake, chocolate, devil's food or fudge, with icing or filling
53105275	Cake or cupcake, chocolate, devil's food or fudge, without icing or filling
53105300	Cake or cupcake, German chocolate, with icing or filling
53105500	Cake, chocolate, with icing, diet
53106500	Cake, cream, without icing or topping
53108200	Snack cake, chocolate, with icing or filling
53108220	Snack cake, chocolate, with icing or filling, reduced fat and calories
53109200	Snack cake, not chocolate, with icing or filling
53109220	Snack cake, not chocolate, with icing or filling, reduced fat and calories
53109300	Cake, Dobos Torte (non-chocolate layer cake with chocolate filling and icing)
53110000	Cake, fruit cake, light or dark, holiday type cake
53111000	Cake or cupcake, gingerbread
53112000	Cake, ice cream and cake roll, chocolate
53112100	Cake, ice cream and cake roll, not chocolate
53113000	Cake, jelly roll
53114000	Cake or cupcake, lemon, without icing or filling
53114100	Cake or cupcake, lemon, with icing or filling
53115100	Cake or cupcake, marble, without icing or filling
53115200	Cake or cupcake, marble, with icing or filling
53115310	Cake or cupcake, nut, without icing or filling
53115320	Cake or cupcake, nut, with icing or filling
53115410	Cake or cupcake, oatmeal
53115450	Cake or cupcake, peanut butter
53116000	Cake, pound, without icing or filling
53116020	Cake, pound, with icing or filling
53116270	Cake, pound, chocolate
53116350	Cake, pound, Puerto Rican style (Ponque)
53116390	Cake, pound, reduced fat, cholesterol free
53116500	Cake or cupcake, pumpkin, without icing or filling
53116510	Cake or cupcake, pumpkin, with icing or filling
53116550	Cake or cupcake, raisin-nut
53116570	Cake, Ravani (made with farina)
53116600	Cake, rice flour, without icing or filling
53116650	Cake, Quezadilla, El Salvadorian style
53117100	Cake or cupcake, spice, without icing or filling
53117200	Cake or cupcake, spice, with icing or filling
53118100	Cake, sponge, without icing or filling
53118200	Cake, sponge, with icing or filling
53118300	Cake, sponge, chocolate
53118410	Rum cake, without icing (Sopa Borracha)
53118500	Cake, torte
53118550	Cake, tres leche
53119000	Cake, upside down (all fruits)
53120270	Cake or cupcake, white, with icing or filling
53120275	Cake or cupcake, white, without icing or filling
53121270	Cake or cupcake, yellow, with icing or filling
53121275	Cake or cupcake, yellow, without icing or filling
53122070	Cake, shortcake, biscuit type, with whipped cream and fruit
53122080	Cake, shortcake, biscuit type, with fruit
53123070	Cake, shortcake, sponge type, with whipped cream and fruit
53123080	Cake, shortcake, sponge type, with fruit
53123500	Cake, shortcake, with whipped topping and fruit, diet

53124110 Cake or cupcake, zucchini  
55801000 Funnel cake with sugar  
55801010 Funnel cake with sugar and fruit

**Cookies****[EPG] = 0.06 (21% adjusted for fat portion of 30%)**

53201000 Cookie, NFS  
53202000 Cookie, almond  
53203000 Cookie, applesauce  
53203500 Cookie, biscotti (Italian sugar cookie)  
53205260 Cookie, bar, with chocolate  
53206000 Cookie, chocolate chip  
53206020 Cookie, chocolate chip, made from home recipe or purchased at a bakery  
53206030 Cookie, chocolate chip, reduced fat  
53206100 Cookie, chocolate chip sandwich  
53206500 Cookie, chocolate, made with rice cereal  
53206550 Cookie, chocolate, made with oatmeal and coconut (no-bake)  
53207000 Cookie, chocolate or fudge  
53207020 Cookie, chocolate or fudge, reduced fat  
53207050 Cookie, chocolate, with chocolate filling or coating, fat free  
53208000 Cookie, marshmallow, chocolate-covered  
53208200 Cookie, marshmallow pie, chocolate covered  
53209005 Cookie, chocolate, with icing or coating  
53209010 Cookie, sugar wafer, chocolate-covered  
53209015 Cookie, chocolate sandwich  
53209020 Cookie, chocolate sandwich, reduced fat  
53209100 Cookie, chocolate, sandwich, with extra filling  
53209500 Cookie, chocolate and vanilla sandwich  
53210000 Cookie, chocolate wafer  
53210900 Cookie, graham cracker with chocolate and marshmallow  
53211000 Cookie bar, with chocolate, nuts, and graham crackers  
53215500 Cookie, coconut  
53220000 Cookie, fruit-filled bar  
53220010 Cookie, fruit-filled bar, fat free  
53220030 Cookie, fig bar  
53220040 Cookie, fig bar, fat free  
53222010 Cookie, fortune  
53222020 Cookie, cone shell, ice cream type, wafer or cake  
53223000 Cookie, gingersnaps  
53223100 Cookie, granola  
53224000 Cookie, ladyfinger  
53224250 Cookie, lemon bar  
53225000 Cookie, macaroon  
53226000 Cookie, marshmallow, with coconut  
53226500 Cookie, marshmallow, with rice cereal (no-bake)  
53226550 Cookie, marshmallow, with rice cereal and chocolate chips  
53226600 Cookie, marshmallow and peanut butter, with oat cereal (no-bake)  
53228000 Cookie, meringue  
53230000 Cookie, molasses  
53231000 Cookie, Lebkuchen  
53231400 Cookie, multigrain, high fiber  
53233000 Cookie, oatmeal  
53233010 Cookie, oatmeal, with raisins  
53233040 Cookie, oatmeal, reduced fat, NS as to raisins



53233050	Cookie, oatmeal sandwich, with creme filling
53233060	Cookie, oatmeal, with chocolate chips
53233080	Cookie, oatmeal sandwich, with peanut butter and jelly filling
53233100	Cookie, oatmeal, with chocolate and peanut butter (no-bake)
53234000	Cookie, peanut butter
53234100	Cookie, peanut butter, with chocolate
53234250	Cookie, peanut butter with rice cereal (no-bake)
53235000	Cookie, peanut butter sandwich
53235500	Cookie, with peanut butter filling, chocolate-coated
53235600	Cookie, Pfeffernusse
53236000	Cookie, pizzelle (Italian style wafer)
53236100	Cookie, pumpkin
53237000	Cookie, raisin
53237010	Cookie, raisin sandwich, cream-filled
53237500	Cookie, rum ball (no-bake)
53238000	Cookie, sandwich-type, not chocolate or vanilla
53239000	Cookie, shortbread
53239010	Cookie, shortbread, reduced fat
53239050	Cookie, shortbread, with icing or filling
53240000	Cookie, animal
53240010	Cookie, animal, with frosting or icing
53241500	Cookie, butter or sugar
53241510	Marie biscuit
53241600	Cookie, butter or sugar, with fruit and/or nuts
53242000	Cookie, sugar wafer
53242500	Cookie, toffee bar
53243000	Cookie, vanilla sandwich
53243010	Cookie, vanilla sandwich, extra filling
53243050	Cookie, vanilla sandwich, reduced fat
53244010	Cookie, butter or sugar, with chocolate icing or filling
53244020	Cookie, butter or sugar, with icing or filling other than chocolate
53246000	Cookie, tea, Japanese
53247000	Cookie, vanilla wafer
53247050	Cookie, vanilla wafer, reduced fat
53247500	Cookie, vanilla with caramel, coconut, and chocolate coating
53251100	Cookie, rugelach
53260030	Cookie, chocolate chip, sugar free
53260200	Cookie, oatmeal, sugar free
53260300	Cookie, sandwich, sugar free
53260400	Cookie, sugar or plain, sugar free
53260500	Cookie, sugar wafer, sugar free
53260600	Cookie, peanut butter, sugar free
53270100	Cookies, Puerto Rican (Mantecaditos polvorones)

**Crepes****[EPG] = 0.06 (20% adjusted for fat portion of 28%)**

53430000	Crepe, dessert type, NS as to filling
53430100	Crepe, dessert type, chocolate-filled
53430200	Crepe, dessert type, fruit-filled
53430250	Crepe suzette
53430300	Crepe, dessert type, ice cream-filled
55401000	Crepe, plain
58120110	Crepes, filled with meat, fish, or poultry, with sauce
58120120	Crepe, filled with beef, pork, fish and/or poultry, no sauce on top

**Desserts excluding cakes, cookies, brownies****[EPG] = 0.06 (20% adjusted for fat portion of 28%)**

14630200	Cheese souffle
27246400	Chicken or turkey souffle
27250550	Seafood souffle
53344200	Mixed fruit tart filled with custard or cream cheese
53344300	Dessert pizza
53391200	Vanilla wafer dessert base
53400200	Blintz, cheese-filled
53400300	Blintz, fruit-filled
53410100	Cobbler, apple
53410200	Cobbler, apricot
53410300	Cobbler, berry
53410500	Cobbler, cherry
53410800	Cobbler, peach
53410850	Cobbler, pear
53410860	Cobbler, pineapple
53410880	Cobbler, plum
53410900	Cobbler, rhubarb
53415100	Crisp, apple, apple dessert
53415120	Fritter, apple
53415200	Fritter, banana
53415220	Fritter, berry
53415300	Crisp, blueberry
53415400	Crisp, cherry
53415500	Crisp, peach
53415600	Crisp, rhubarb
53420000	Cream puff, eclair, custard or cream filled, NS as to icing
53420100	Cream puff, eclair, custard or cream filled, not iced
53420200	Cream puff, eclair, custard or cream filled, iced
53420210	Cream puff, eclair, custard or cream filled, iced, reduced fat
53420250	Cream puff, no filling or icing
53420300	Air filled fritter or fried puff, without syrup, Puerto Rican style (Bunuelos de viento)
53420310	Wheat flour fritter, without syrup
53420400	Sopaipilla, without syrup or honey
53420410	Sopaipilla with syrup or honey
53430700	Tamale, sweet
53430750	Tamale, sweet, with fruit
53441110	Baklava
53441210	Basbousa (semolina dessert dish)
63403150	Lime souffle
72125240	Spinach souffle
73305020	Squash, winter, souffle
75418060	Squash, summer, soufflé

**Doughnuts****[EPG] = 0.03% (12% adjusted for fat portion of 23%)**

53520000	Doughnut, NS as to cake or yeast
53520110	Doughnut, cake type
53520120	Doughnut, cake type, chocolate
53520140	Doughnut, cake type, chocolate covered
53520150	Doughnut, cake type, chocolate covered, dipped in peanuts
53520160	Doughnut, cake type, chocolate, with chocolate icing

53520500	Doughnut, Asian
53520600	Cruller, NFS
53520700	French cruller
53521100	Doughnut, raised or yeast, chocolate, with chocolate icing
53521110	Doughnut, raised or yeast
53521120	Doughnut, raised or yeast, chocolate
53521130	Doughnut, raised or yeast, chocolate covered
53521140	Doughnut, jelly
53521210	Doughnut, custard-filled
53521220	Doughnut, chocolate cream-filled
53521230	Doughnut, custard-filled, with icing
53521250	Doughnut, wheat
53521300	Doughnut, wheat, chocolate covered

**Muffins****[EPG] = 0.03% (13% adjusted for fat portion of 19%)**

52301000	Muffin, NFS
52302010	Muffin, fruit and/or nuts
52302020	Muffin, fruit, low fat
52302500	Muffin, chocolate chip
52302600	Muffin, chocolate
52303010	Muffin, whole wheat
52303500	Muffin, wheat
52304000	Muffin, whole grain
52304010	Muffin, wheat bran
52304040	Muffin, bran with fruit, lowfat
52304100	Muffin, oatmeal
52304150	Muffin, oat bran
52304200	Muffin, oat bran with fruit and/or nuts
52306010	Muffin, plain
52306300	Muffin, cheese
52306500	Muffin, pumpkin
52306550	Muffin, zucchini
52306700	Muffin, carrot

**Pastry****[EPG] = 0.03% (15% adjusted for fat portion of 21%)**

53440000	Strudel, apple
53440300	Strudel, berry
53440500	Strudel, cherry
53440600	Strudel, cheese
53440700	Strudel, peach
53440750	Strudel, pineapple
53440800	Strudel, cheese and fruit
53450000	Turnover or dumpling, apple
53450300	Turnover or dumpling, berry
53450500	Turnover or dumpling, cherry
53450800	Turnover or dumpling, lemon
53451000	Turnover or dumpling, peach
53451500	Turnover, guava
53451750	Turnover, pumpkin
53452100	Pastry, fruit-filled
53452120	Pastry, Asian, made with bean or lotus seed paste filling (baked)

53452130	Pastry, Asian, made with bean paste and salted egg yolk filling (baked)
53452150	Pastry, Chinese, made with rice flour
53452170	Pastry, cookie type, fried
53452200	Pastry, Italian, with cheese
53452400	Pastry, puff
53452420	Pastry, puff, custard or cream filled, iced or not iced
53452450	Cheese pastry puffs
53452500	Pastry, mainly flour and water, fried
53453150	Empanada, Mexican turnover, fruit-filled
53453170	Empanada, Mexican turnover, pumpkin
52105100	Scone
52105110	Scone, whole wheat
52105200	Scone, with fruit

## Pastry crusts and pies

[EPG] = 0.09% (25% adjusted for fat portion of 36%)

53300100	Pie, NFS
53300170	Pie, individual size or tart, NFS
53300180	Pie, fried, NFS
53301000	Pie, apple, two crust
53301070	Pie, apple, individual size or tart
53301080	Pie, apple, fried pie
53301500	Pie, apple, one crust
53301750	Pie, apple, diet
53302000	Pie, apricot, two crust
53302070	Pie, apricot, individual size or tart
53302080	Pie, apricot, fried pie
53303000	Pie, blackberry, two crust
53303070	Pie, blackberry, individual size or tart
53303500	Pie, berry, not blackberry, blueberry, boysenberry, huckleberry, raspberry, or strawberry; two crust
53303510	Pie, berry, not blackberry, blueberry, boysenberry, huckleberry, raspberry, or strawberry; one crust
53303570	Pie, berry, not blackberry, blueberry, boysenberry, huckleberry, raspberry, or strawberry, individual size or tart
53304000	Pie, blueberry, two crust
53304050	Pie, blueberry, one crust
53304070	Pie, blueberry, individual size or tart
53305000	Pie, cherry, two crust
53305010	Pie, cherry, one crust
53305070	Pie, cherry, individual size or tart
53305080	Pie, cherry, fried pie
53305700	Pie, lemon (not cream or meringue)
53305720	Pie, lemon (not cream or meringue), individual size or tart
53305750	Pie, lemon, fried pie
53306000	Pie, mince, two crust
53306070	Pie, mince, individual size or tart
53307000	Pie, peach, two crust
53307050	Pie, peach, one crust
53307070	Pie, peach, individual size or tart
53307080	Pie, peach, fried pie
53307500	Pie, pear, two crust
53307570	Pie, pear, individual size or tart
53308000	Pie, pineapple, two crust

53308070	Pie, pineapple, individual size or tart
53308300	Pie, plum, two crust
53308500	Pie, prune, one crust
53309000	Pie, raisin, two crust
53309070	Pie, raisin, individual size or tart
53310000	Pie, raspberry, one crust
53310050	Pie, raspberry, two crust
53311000	Pie, rhubarb, two crust
53311050	Pie, rhubarb, one crust
53311070	Pie, rhubarb, individual size or tart
53312000	Pie, strawberry, one crust
53313000	Pie, strawberry-rhubarb, two crust
53314000	Pie, strawberry, individual size or tart
53340000	Pie, apple-sour cream
53340500	Pie, cherry, made with cream cheese and sour cream
53341000	Pie, banana cream
53341070	Pie, banana cream, individual size or tart
53341500	Pie, buttermilk
53341750	Pie, chess
53342000	Pie, chocolate cream
53342070	Pie, chocolate cream, individual size or tart
53343000	Pie, coconut cream
53343070	Pie, coconut cream, individual size or tart
53344000	Pie, custard
53344070	Pie, custard, individual size or tart
53345000	Pie, lemon cream
53345070	Pie, lemon cream, individual size or tart
53346000	Pie, peanut butter cream
53346500	Pie, pineapple cream
53347000	Pie, pumpkin
53347070	Pie, pumpkin, individual size or tart
53347100	Pie, raspberry cream
53347500	Pie, sour cream, raisin
53347600	Pie, squash
53348000	Pie, strawberry cream
53348070	Pie, strawberry cream, individual size or tart
53360000	Pie, sweet potato
53365000	Pie, vanilla cream
53366000	Pie, yogurt, frozen
53370000	Pie, chiffon, not chocolate
53371000	Pie, chiffon, chocolate
53371100	Pie, chiffon, with liqueur
53373000	Pie, black bottom
53381000	Pie, lemon meringue
53381070	Pie, lemon meringue, individual size or tart
53382000	Pie, chocolate-marshmallow
53385000	Pie, pecan
53385070	Pie, pecan, individual size or tart
53385500	Pie, oatmeal
53386000	Pie, pudding, flavors other than chocolate
53386050	Pie, pudding, flavors other than chocolate, individual size or tart
53386250	Pie, pudding, chocolate, with chocolate coating, individual size
53386500	Pie, pudding, flavors other than chocolate, with chocolate coating, individual size
53387000	Pie, Toll house chocolate chip
53390000	Pie, shoo-fly

53390100	Pie, tofu with fruit
53391000	Pie shell
53391100	Pie shell, graham cracker
53391150	Pie shell, chocolate wafer

**Frozen Dairy Desserts and Mixes****Ice Cream****[EPG] = 0.03% (12% adjusted for fat portion of 21%)**

13110000	Ice cream, NFS
13110100	Ice cream, regular, flavors other than chocolate
13110110	Ice cream, regular, chocolate
13110120	Ice cream, rich, flavors other than chocolate
13110130	Ice cream, rich, chocolate
13110140	Ice cream, rich, NS as to flavor
13110200	Ice cream, soft serve, flavors other than chocolate
13110210	Ice cream, soft serve, chocolate
13110220	Ice cream, soft serve, NS as to flavor
13110310	Ice cream, no sugar added, NS as to flavor
13110320	Ice cream, no sugar added, flavors other than chocolate
13110330	Ice cream, no sugar added, chocolate
13120050	Ice cream bar or stick, not chocolate covered or cake covered
13120310	Ice cream bar, stick or nugget, with crunch coating
13120740	Ice cream cone, no topping, NS as to flavor
13121200	Ice cream sundae, prepackaged type, flavors other than chocolate
13130100	Light ice cream, NFS (formerly ice milk)
13130300	Light ice cream, flavors other than chocolate (formerly ice milk)
13130310	Light ice cream, chocolate (formerly ice milk)
13130320	Light ice cream, no sugar added, NS as to flavor
13130330	Light ice cream, no sugar added, flavors other than chocolate
13130340	Light ice cream, no sugar added, chocolate
13130590	Light ice cream, soft serve, NS as to flavor (formerly ice milk)
13130600	Light ice cream, soft serve, flavors other than chocolate (formerly ice milk)
13130610	Light ice cream, soft serve, chocolate (formerly ice milk)
13130640	Light ice cream, soft serve cone, NS as to flavor (formerly ice milk)
13130700	Light ice cream, soft serve, blended with candy or cookies
13135010	Ice cream sandwich, made with light chocolate ice cream
13136000	Ice cream sandwich, made with light, no sugar added ice cream
13140570	Light ice cream, no sugar added, cone, NS as to flavor
13140575	Light ice cream, no sugar added, cone, flavors other than chocolate
13140580	Light ice cream, no sugar added, cone, chocolate
13140710	Light ice cream, creamsicle or dreamsicle, no sugar added
13160150	Fat free ice cream, no sugar added, chocolate
13160160	Fat free ice cream, no sugar added, flavors other than chocolate
13160400	Fat free ice cream, flavors other than chocolate
13160410	Fat free ice cream, chocolate
13160420	Fat free ice cream, NS as to flavor
13161630	Light ice cream, bar or stick, with low-calorie sweetener, chocolate-coated (formerly ice milk)

Mixed foods containing ice cream*Adjusted for ice cream content of ~6 to 11.52%*

13120100	Ice cream bar or stick, chocolate covered
13120110	Ice cream bar or stick, chocolate or caramel covered, with nuts
13120120	Ice cream bar or stick, rich chocolate ice cream, thick chocolate covering
13120121	Ice cream bar or stick, rich ice cream, thick chocolate covering
13120130	Ice cream bar or stick, rich ice cream, chocolate covered, with nuts
13120140	Ice cream bar or stick, chocolate ice cream, chocolate covered
13120400	Ice cream bar or stick with fruit
13120500	Ice cream sandwich
13120550	Ice cream cookie sandwich
13120700	Ice cream cone with nuts, flavors other than chocolate
13120710	Ice cream cone, chocolate covered, with nuts, flavors other than chocolate
13120720	Ice cream cone, chocolate covered or dipped, flavors other than chocolate
13120730	Ice cream cone, no topping, flavors other than chocolate
13120750	Ice cream cone with nuts, chocolate ice cream
13120760	Ice cream cone, chocolate covered or dipped, chocolate ice cream
13120770	Ice cream cone, no topping, chocolate ice cream
13120780	Ice cream cone, chocolate covered, with nuts, chocolate ice cream
13120790	Ice cream sundae cone
13121000	Ice cream sundae, NS as to topping, with whipped cream
13121100	Ice cream sundae, fruit topping, with whipped cream
13121300	Ice cream sundae, chocolate or fudge topping, with whipped cream
13121400	Ice cream sundae, not fruit or chocolate topping, with whipped cream
13122100	Ice cream pie, no crust
13126000	Ice cream, fried
13130620	Light ice cream, soft serve cone, flavors other than chocolate (formerly ice milk)
13130630	Light ice cream, soft serve cone, chocolate (formerly ice milk)
13135000	Light ice cream, sandwich (formerly ice milk)
13140100	Light ice cream, bar or stick, chocolate-coated (formerly ice milk)
13140110	Light ice cream, bar or stick, chocolate covered, with nuts (formerly ice milk)
13140500	Light ice cream, cone, flavors other than chocolate (formerly ice milk)
13140550	Light ice cream, cone, chocolate (formerly ice milk)
13140600	Light ice cream, sundae, soft serve, chocolate or fudge topping, with whipped cream (formerly ice milk)
13140630	Light ice cream, sundae, soft serve, fruit topping, with whipped cream (formerly ice milk)
13140660	Light ice cream, sundae, soft serve, chocolate or fudge topping (without whipped cream)
13140670	Light ice cream, sundae, soft serve, fruit topping (without whipped cream) (formerly ice milk)
13140680	Light ice cream, sundae, soft serve, not fruit or chocolate topping (without whipped cream)
13140900	Light ice cream, fudgesicle (formerly ice milk)

Mixed foods containing ice cream*Adjusted for ice cream content of ~4 to ~49%*

11541000	Milk shake, NS as to flavor or type
11541100	Milk shake, homemade or fountain-type, NS as to flavor
11541110	Milk shake, homemade or fountain-type, chocolate
11541120	Milk shake, homemade or fountain-type, flavors other than chocolate
11541400	Milk shake with malt
13120300	Ice cream bar, cake covered
13120800	Ice cream soda, flavors other than chocolate
13120810	Ice cream soda, chocolate

13121500	Ice cream sundae, fudge topping, with cake, with whipped cream
13122500	Ice cream pie, with cookie crust, fudge topping, and whipped cream
13140450	Light ice cream, cone, NFS (formerly ice milk)
13140650	Light ice cream, sundae, soft serve, not fruit or chocolate topping, with whipped cream
13140700	Light ice cream, creamsicle or dreamsicle (formerly ice milk)
13170000	Baked Alaska
91611050	Ice pop filled with ice cream, all flavor varieties

## Grain Products and Pastas

### Granola Bars

[EPG] = 0.04% (17% adjusted for fat portion of 24%)

53710400	Fiber One Chewy Bar
53710500	Kellogg's Nutri-Grain Cereal Bar
53710502	Kellogg's Nutri-Grain Yogurt Bar
53710504	Kellogg's Nutri-Grain Fruit and Nut Bar
53710600	Milk 'n Cereal bar
53710700	Kellogg's Special K bar
53710800	Kashi GOLEAN Chewy Bars
53710802	Kashi TLC Chewy Granola Bar
53710804	Kashi GOLEAN Crunchy Bars
53710806	Kashi TLC Crunchy Granola Bar
53710900	Nature Valley Chewy Trail Mix Granola Bar
53710902	Nature Valley Chewy Granola Bar with Yogurt Coating
53710904	Nature Valley Sweet and Salty Granola Bar
53710906	Nature Valley Crunchy Granola Bar
53711000	Quaker Chewy Granola Bar
53711002	Quaker Chewy 90 Calorie Granola Bar
53711004	Quaker Chewy 25% Less Sugar Granola Bar
53711006	Quaker Chewy Dipps Granola Bar
53711100	Quaker Granola Bites
53712000	Snack bar, oatmeal
53712100	Granola bar, NFS
53712200	Granola bar, lowfat, NFS
53712210	Granola bar, nonfat
53713000	Granola bar, reduced sugar, NFS
53713100	Granola bar, peanuts , oats, sugar, wheat germ
53714200	Granola bar, chocolate-coated, NFS
53714210	Granola bar, with coconut, chocolate-coated
53714220	Granola bar with nuts, chocolate-coated
53714230	Granola bar, oats, nuts, coated with non-chocolate coating
53714250	Granola bar, coated with non-chocolate coating
53714300	Granola bar, high fiber, coated with non-chocolate yogurt coating
53714400	Granola bar, with rice cereal
53714500	Breakfast bar, NFS
53714510	Breakfast bar, date, with yogurt coating
53714520	Breakfast bar, cereal crust with fruit filling, lowfat
53720100	Balance Original Bar
53720200	Clif Bar
53720300	PowerBar
53720400	Slim Fast Original Meal Bar
53720500	Snickers Marathon Protein bar
53720510	Snickers Marathon Energy bar



53720600	South Beach Living Meal Bar
53720610	South Beach Living High Protein Bar
53720700	Tiger's Milk bar
53720800	Zone Perfect Classic Crunch nutrition bar
53729000	Nutrition bar or meal replacement bar, NFS

**Gravies and Sauces****Pasta sauces (cream and tomato based)****[EPG] = 0.01% (8% adjusted for fat portion of 11%)**

14650160	Alfredo sauce
74403010	Tomato sauce
74403050	Tomato sauce, low sodium
74404010	Spaghetti sauce
74404060	Spaghetti sauce, fat free
74404090	Vodka flavored pasta sauce made with tomatoes and cream

**Mixed foods containing pasta sauces (cream and tomato based)*****Adjusted for pasta sauces content of ~54 to 94%***

27120110	Sausage with tomato-based sauce (mixture)
58132110	Spaghetti with tomato sauce, meatless
58132350	Spaghetti with tomato sauce, meatless, whole wheat noodles
58132450	Spaghetti with tomato sauce, meatless, made with spinach noodles
58146110	Pasta with meat sauce
74404030	Spaghetti sauce with meat, canned, no extra meat added
75306010	Eggplant in tomato sauce, cooked, fat not added in cooking

**Mixed foods containing pasta sauces (cream and tomato based)*****Adjusted for pasta sauces content of ~25 to ~48% and fat portion of 11%***

27111050	Spaghetti sauce with beef or meat other than lamb or mutton, homemade-style
27111500	Beef sloppy joe (no bun)
27120100	Ham or pork with tomato-based sauce (mixture)
27120250	Frankfurters or hot dogs with tomato-based sauce (mixture)
27130040	Spaghetti sauce with lamb or mutton, homemade-style
27135110	Veal parmigiana
27136050	Venison/deer with tomato-based sauce (mixture)
27146150	Chicken curry
27146300	Chicken or turkey parmigiana
27212100	Beef and noodles with tomato-based sauce (mixture)
27242400	Chicken or turkey and noodles, tomato-based sauce (mixture)
27319010	Stuffed green pepper, Puerto Rican style (Pimiento relleno)
27320100	Pork, potatoes, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based
27320110	Pork, potatoes, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based sau
27320340	Pork, rice, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato- based sauc
27320350	Pork, rice, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato- based sauce (
27411200	Beef with vegetables (excluding carrots, broccoli, and dark-green leafy (no potatoes)), tomato-based

28140740	Chicken patty, or nuggets, boneless, breaded, with pasta and tomato sauce, fruit, dessert (frozen me
32110100	Eggs a la Malaguena, Puerto Rican style (Huevos a la Malaguena)
58103210	Tamale, meatless, Puerto Rican or Caribbean style
58109000	Italian pie, meatless
58109010	Italian pie with meat
58131110	Ravioli, NS as to filling, with tomato sauce
58131520	Ravioli, cheese-filled, with tomato sauce
58131530	Ravioli, cheese-filled, with meat sauce
58132310	Spaghetti with tomato sauce and meatballs or spaghetti with meat sauce or spaghetti with meat sauce and meatballs
58132360	Spaghetti with tomato sauce and meatballs, whole wheat noodles or spaghetti with meat sauce, whole wheat noodles or spaghetti with meat sauce and meatballs, whole wheat noodles
58132460	Spaghetti with tomato sauce and meatballs made with spinach noodles, or spaghetti with meat sauce made with spinach noodles, or spaghetti with meat sauce and meatballs made with spinach noodles
58132710	Spaghetti with tomato sauce and frankfurters or hot dogs
58132910	Spaghetti with tomato sauce and chicken or turkey
58133120	Manicotti, cheese-filled, with tomato sauce, meatless
58133140	Manicotti, vegetable- and cheese-filled, with tomato sauce, meatless
58134120	Stuffed shells, cheese-filled, with tomato sauce, meatless
58134620	Tortellini, cheese-filled, meatless, with tomato sauce
58146100	Pasta with tomato sauce, meatless
58146120	Pasta with cheese and meat sauce
58146130	Pasta with carbonara sauce
58146150	Pasta with cheese and tomato sauce, meatless
58146160	Pasta with vegetables, no sauce or dressing
58147100	Pasta with pesto sauce
58161300	White rice with tomato sauce
58161310	Rice, brown, with tomato sauce
58421000	Sopa seca (dry soup), Mexican style, NFS
58421010	Sopa Seca de Fideo, Mexican style, made with dry noodles
58421060	Sopa seca de arroz (dry rice soup), Mexican style
75302010	Beans, green string, with tomatoes, cooked, fat not added in cooking
75412070	Eggplant with cheese and tomato sauce
75418030	Squash, summer, casserole, with rice and tomato sauce

Mixed foods containing pasta sauces (cream and tomato based)

*Adjusted for pasta sauces content of ~2 to ~<25%*

58132710	Spaghetti with tomato sauce and frankfurters or hot dogs
58132910	Spaghetti with tomato sauce and chicken or turkey
58146100	Pasta with tomato sauce, meatless
58146130	Pasta with carbonara sauce
58146150	Pasta with cheese and tomato sauce, meatless
58146160	Pasta with vegetables, no sauce or dressing
58147100	Pasta with pesto sauce
27130040	Spaghetti sauce with lamb or mutton, homemade-style
27111050	Spaghetti sauce with beef or meat other than lamb or mutton, homemade-style
27120250	Frankfurters or hot dogs with tomato-based sauce (mixture)
27411200	Beef with vegetables (excluding carrots, broccoli, and dark-green leafy (no potatoes)), tomato-based
27120100	Ham or pork with tomato-based sauce (mixture)

27320350	Pork, rice, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based sauce
27212100	Beef and noodles with tomato-based sauce (mixture)
58132310	Spaghetti with tomato sauce and meatballs or spaghetti with meat sauce or spaghetti with meat sauce and meatballs
58132360	Spaghetti with tomato sauce and meatballs, whole wheat noodles or spaghetti with meat sauce, whole wheat noodles or spaghetti with meat sauce and meatballs, whole wheat noodles
58132460	Spaghetti with tomato sauce and meatballs made with spinach noodles, or spaghetti with meat sauce made with spinach noodles, or spaghetti with meat sauce and meatballs made with spinach noodles
28140740	Chicken patty, or nuggets, boneless, breaded, with pasta and tomato sauce, fruit, dessert (frozen meat)
27242400	Chicken or turkey and noodles, tomato-based sauce (mixture)
27320340	Pork, rice, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based sauce
75302010	Beans, green string, with tomatoes, cooked, fat not added in cooking
27320100	Pork, potatoes, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based
27320110	Pork, potatoes, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based sauce
58161300	White rice with tomato sauce
75418030	Squash, summer, casserole, with rice and tomato sauce
58421010	Sopa Seca de Fideo, Mexican style, made with dry noodles
58134620	Tortellini, cheese-filled, meatless, with tomato sauce
27136050	Venison/deer with tomato-based sauce (mixture)
32110100	Eggs a la Malaguena, Puerto Rican style (Huevos a la Malaguena)
58131520	Ravioli, cheese-filled, with tomato sauce
58421000	Sopa seca (dry soup), Mexican style, NFS
58131110	Ravioli, NS as to filling, with tomato sauce
58421060	Sopa seca de arroz (dry rice soup), Mexican style
58131530	Ravioli, cheese-filled, with meat sauce
58161310	Rice, brown, with tomato sauce
58103210	Tamale, meatless, Puerto Rican or Caribbean style
58134120	Stuffed shells, cheese-filled, with tomato sauce, meatless
58109000	Italian pie, meatless
27111500	Beef sloppy joe (no bun)
27135110	Veal parmigiana
27146300	Chicken or turkey parmigiana
58146120	Pasta with cheese and meat sauce
58109010	Italian pie with meat
58133140	Manicotti, vegetable- and cheese-filled, with tomato sauce, meatless
58133120	Manicotti, cheese-filled, with tomato sauce, meatless
27319010	Stuffed green pepper, Puerto Rican style (Pimiento relleno)
27146150	Chicken curry
75412070	Eggplant with cheese and tomato sauce
75412060	Eggplant parmesan casserole, regular
27510700	Meatball and spaghetti sauce submarine sandwich, on roll
58133130	Manicotti, cheese-filled, with meat sauce
27320080	Sausage, noodles, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based
58146310	Pasta, whole wheat, with tomato sauce, meatless
27560720	Sausage and spaghetti sauce sandwich
27320090	Sausage, noodles, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based

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58145140	Macaroni or noodles with cheese and tomato
28510010	Gravy or sauce, poultry-based from Puerto Rican-style chicken fricasse
27220110	Pork and rice with tomato-based sauce (mixture)
27118110	Meatballs, Puerto Rican style (Albondigas)
27451030	Lobster creole, Puerto Rican style (Langosta a la criolla)
58146300	Pasta, whole wheat, with meat sauce
27510710	Pizzaburger (hamburger, cheese, sauce) on 1/2 bun
77316010	Stuffed cabbage, with meat, Puerto Rican style (Repollo relleno con carne)
75410500	Chiles rellenos, cheese-filled (stuffed chili peppers)
58134610	Tortellini, meat-filled, with tomato sauce
58134710	Tortellini, spinach-filled, with tomato sauce
27330220	Lamb or mutton stew with potatoes and vegetables (excluding carrots, broccoli, and dark-green leafy)
27330210	Lamb or mutton stew with potatoes and vegetables (including carrots, broccoli, and/or dark-green leafy)
27336310	Venison/deer, noodles, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based
27450700	Fish and vegetables (including carrots, broccoli, and/or dark-green leafy (no potatoes)), tomato-based
27336300	Venison/deer, noodles, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato
27450710	Fish and vegetables (excluding carrots, broccoli, and dark-green leafy (no potatoes)), tomato-based
27111420	Chili con carne without beans
27214110	Meat loaf made with beef, with tomato-based sauce
27260100	Meat loaf made with beef and pork, with tomato-based sauce
27510720	Pizzaburger (hamburger, cheese, sauce) on whole bun
27315250	Stuffed cabbage rolls with beef and rice
58106780	Pizza with meat and vegetables, lowfat, thin crust
27111300	Mexican style beef stew, no potatoes, tomato-based sauce (mixture) (Carne guisada sin papas)
27120130	Mexican style pork stew, no potatoes, tomato-based sauce (mixture) (cerdo guisado sin papas)
27141500	Chili con carne with chicken or turkey and beans
58101830	Mexican casserole made with ground beef, tomato sauce, cheese, taco seasonings, and corn chips
27118120	Stewed seasoned ground beef, Puerto Rican style (Picadillo guisado, picadillo de carne)
27141050	Stewed chicken with tomato-based sauce, Mexican style (mixture) (Pollo guisado con tomate)
27136100	Chili con carne with venison/deer and beans
27111400	Chili con carne, NS as to beans
27111410	Chili con carne with beans
27121410	Chili con carne with beans, made with pork
58100240	Burrito with chicken, NFS
27461010	Stewed seasoned ground beef, Puerto Rican style (Picadillo para relleno)
27441120	Chicken or turkey creole, without rice
27211110	Mexican style beef stew with potatoes, tomato-based sauce (mixture) (Carne guisada con papas)
27221150	Mexican style pork stew, with potatoes, tomato-based sauce (mixture) (cerdo guisado con papas)
27221110	Pork roast, stuffed, Puerto Rican style
58126150	Turnover, meat- and cheese-filled, tomato-based sauce
58108030	Panzerotti, with meat, vegetables, and cheese
58100800	Enchilada with cheese, meatless, no beans
58100530	Enchilada with beef and cheese, no beans

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27515050	Fajita-style beef sandwich with cheese, on pita bread, with lettuce and tomato
27540200	Fajita-style chicken sandwich with cheese, on pita bread, with lettuce and tomato
27451010	Fish a la creole, Puerto Rican style (Pescado frito con mojo)
58100630	Enchilada with chicken and cheese, no beans, tomato-based sauce
58101820	Mexican casserole made with ground beef, beans, tomato sauce, cheese, taco seasonings, and corn chip
58100400	Enchilada with beef, no beans
27418310	Corned beef with tomato sauce and onion, Puerto Rican style (mixture)
27111310	Mexican style beef stew, no potatoes, with chili peppers, tomato-based sauce (mixture) (Carne guisada con papas)
58100560	Enchilada with ham and cheese, no beans
27212120	Chili con carne with beans and macaroni
58100900	Enchilada with seafood, tomato-based sauce
58100520	Enchilada with beef, beans, and cheese
71931010	Cassava with creole sauce, Puerto Rican style (Yuca al mojo)
58100620	Enchilada with chicken, beans, and cheese, tomato-based sauce
58100600	Enchilada with chicken, tomato-based sauce
58100230	Burrito with chicken and cheese
27118130	Stewed dried beef, Puerto Rican style (Tasajo guisado, carne cecina guisada)
58100720	Enchilada with beans and cheese, meatless
58100610	Enchilada with chicken and beans, tomato-based sauce
27330060	Lamb or mutton, rice, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato
58100510	Enchilada with beef and beans
77141010	Potato chicken pie, Puerto Rican style (Pastelon de pollo)
41310210	Stewed chickpeas with Spanish sausages, Puerto Rican style (Garbanzos guisados con chorizos)
28510030	Gravy, meat-based, from Puerto-Rican style beef stew
27260500	Vienna sausages stewed with potatoes, Puerto Rican style (Salchichas guisadas)
27351040	Biscayne codfish, Puerto Rican style (Bacalao a la Vizcaina)
58100710	Enchilada with beans, meatless
58140110	Spaghetti with corned beef, Puerto Rican style
58100220	Burrito with chicken, beans, and cheese
77316510	Stuffed cabbage, with meat and rice, Syrian dish, Puerto Rican style
27218310	Stewed corned beef, Puerto Rican style ("Corned beef" guisado)
27331150	Veal fricassee, Puerto Rican style (ternera en fricase)
74415110	Tomato and sofrito stewing sauce, Puerto Rican style
27335500	Stewed rabbit, Puerto Rican style (Fricase de conejo)
77250110	Stuffed tannier fritters, Puerto Rican style (Alcapurrias)
27348100	Chicken fricassee, Puerto Rican style (Fricase de pollo)
27420410	Pork and vegetables (excluding carrots, broccoli, and dark-green leafy (no potatoes)), tomato-based
41210100	Stewed dry red beans, Puerto Rican style (Habichuelas coloradas guisadas)
27420400	Pork and vegetables (including carrots, broccoli, and/or dark-green leafy (no potatoes)), tomato-based
27462000	Stewed chitterlings, Puerto Rican style (cuajo guisado)
58155810	Stewed rice, Puerto Rican style (arroz quisado)
27118180	Puerto Rican style beef stew, meat with gravy (potatoes reported separately)
77205110	Ripe plantain fritters, Puerto Rican style (Pionono)
27218210	Puerto-Rican style beef stew (Carne guisada con papas)
58156210	Rice with vienna sausage, Puerto Rican style (arroz con salchichas)
58156310	Rice with Spanish sausage, Puerto Rican style
58155210	Stuffed rice with chicken, Dominican style (Arroz relleno Dominicano)
58108040	Panzerotti, with vegetables and cheese
58155910	Rice with squid, Puerto Rican style (arroz con calamares)

58155110	Rice with chicken, Puerto Rican style (Arroz con Pollo)
41310200	Chickpeas stewed with pig's feet, Puerto Rican style (Garbanzos guisados con patitos de cerdo)
77230510	Cassava pie stuffed with crab meat, Puerto Rican style (Empanada de jueyes)
28315140	Beef vegetable soup, Mexican style (Sopa / caldo de Res)
77563010	Puerto Rican stew (Sancocho)
27111300	Mexican style beef stew, no potatoes, tomato-based sauce (mixture) (Carne guisada sin papas)
27111310	Mexican style beef stew, no potatoes, with chili peppers, tomato-based sauce (mixture) (Carne guisad
27111400	Chili con carne, NS as to beans
27111410	Chili con carne with beans
27111420	Chili con carne without beans
27118110	Meatballs, Puerto Rican style (Albondigas)
27118120	Stewed seasoned ground beef, Puerto Rican style (Picadillo guisado, picadillo de carne)
27118130	Stewed dried beef, Puerto Rican style (Tasajo guisado, carne cecina guisada)
27118180	Puerto Rican style beef stew, meat with gravy (potatoes reported separately)
27120130	Mexican style pork stew, no potatoes, tomato-based sauce (mixture) (cerdo guisado sin papas)
27121410	Chili con carne with beans, made with pork
27136100	Chili con carne with venison/deer and beans
27141050	Stewed chicken with tomato-based sauce, Mexican style (mixture) (Pollo guisado con tomate)
27141500	Chili con carne with chicken or turkey and beans
27211110	Mexican style beef stew with potatoes, tomato-based sauce (mixture) (Carne guisada con papas)
27212120	Chili con carne with beans and macaroni
27214110	Meat loaf made with beef, with tomato-based sauce
27218210	Puerto-Rican style beef stew (Carne guisada con papas)
27218310	Stewed corned beef, Puerto Rican style ("Corned beef" guisado)
27220110	Pork and rice with tomato-based sauce (mixture)
27221110	Pork roast, stuffed, Puerto Rican style
27221150	Mexican style pork stew, with potatoes, tomato-based sauce (mixture) (cerdo guisado con papas)
27260100	Meat loaf made with beef and pork, with tomato-based sauce
27260500	Vienna sausages stewed with potatoes, Puerto Rican style (Salchichas guisadas)
27315250	Stuffed cabbage rolls with beef and rice
27320080	Sausage, noodles, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based
27320090	Sausage, noodles, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based
27330060	Lamb or mutton, rice, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato-based
27330210	Lamb or mutton stew with potatoes and vegetables (including carrots, broccoli, and/or dark-green leafy)
27330220	Lamb or mutton stew with potatoes and vegetables (excluding carrots, broccoli, and dark-green leafy)
27331150	Veal fricassee, Puerto Rican style (ternera en fricase)
27335500	Stewed rabbit, Puerto Rican style (Fricase de conejo)
27336300	Venison/deer, noodles, and vegetables (including carrots, broccoli, and/or dark-green leafy), tomato
27336310	Venison/deer, noodles, and vegetables (excluding carrots, broccoli, and dark-green leafy), tomato-based
27348100	Chicken fricassee, Puerto Rican style (Fricase de pollo)
27351040	Biscayne codfish, Puerto Rican style (Bacalao a la Vizcaina)

# Intertek

27418310	Corned beef with tomato sauce and onion, Puerto Rican style (mixture)
27420400	Pork and vegetables (including carrots, broccoli, and/or dark-green leafy (no potatoes)), tomato-based
27420410	Pork and vegetables (excluding carrots, broccoli, and dark- green leafy (no potatoes)), tomato-based
27441120	Chicken or turkey creole, without rice
27450700	Fish and vegetables (including carrots, broccoli, and/or dark-green leafy (no potatoes)), tomato-based
27450710	Fish and vegetables (excluding carrots, broccoli, and dark- green leafy (no potatoes)), tomato-based
27451010	Fish a la creole, Puerto Rican style (Pescado frito con mojo)
27451030	Lobster creole, Puerto Rican style (Langosta a la criolla)
27461010	Stewed seasoned ground beef, Puerto Rican style (Picadillo para relleno)
27462000	Stewed chitterlings, Puerto Rican style (cuajo guisado)
27510700	Meatball and spaghetti sauce submarine sandwich, on roll
27510710	Pizzaburger (hamburger, cheese, sauce) on 1/2 bun
27510720	Pizzaburger (hamburger, cheese, sauce) on whole bun
27515050	Fajita-style beef sandwich with cheese, on pita bread, with lettuce and tomato
27540200	Fajita-style chicken sandwich with cheese, on pita bread, with lettuce and tomato
27560720	Sausage and spaghetti sauce sandwich
28315140	Beef vegetable soup, Mexican style (Sopa / caldo de Res)
28510010	Gravy or sauce, poultry-based from Puerto Rican-style chicken fricasse
28510030	Gravy, meat-based, from Puerto-Rican style beef stew
41210100	Stewed dry red beans, Puerto Rican style (Habichuelas coloradas guisadas)
41310200	Chickpeas stewed with pig's feet, Puerto Rican style (Garbanzos guisados con patitos de cerdo)
41310210	Stewed chickpeas with Spanish sausages, Puerto Rican style (Garbanzos guisados con chorizos)
58100220	Burrito with chicken, beans, and cheese
58100230	Burrito with chicken and cheese
58100240	Burrito with chicken, NFS
58100400	Enchilada with beef, no beans
58100510	Enchilada with beef and beans
58100520	Enchilada with beef, beans, and cheese
58100530	Enchilada with beef and cheese, no beans
58100560	Enchilada with ham and cheese, no beans
58100600	Enchilada with chicken, tomato-based sauce
58100610	Enchilada with chicken and beans, tomato-based sauce
58100620	Enchilada with chicken, beans, and cheese, tomato- based sauce
58100630	Enchilada with chicken and cheese, no beans, tomato- based sauce
58100710	Enchilada with beans, meatless
58100720	Enchilada with beans and cheese, meatless
58100800	Enchilada with cheese, meatless, no beans
58100900	Enchilada with seafood, tomato-based sauce
58101820	Mexican casserole made with ground beef, beans, tomato sauce, cheese, taco seasonings, and corn chip
58101830	Mexican casserole made with ground beef, tomato sauce, cheese, taco seasonings, and corn chips
58106780	Pizza with meat and vegetables, lowfat, thin crust
58108030	Panzerotti, with meat, vegetables, and cheese
58108040	Panzerotti, with vegetables and cheese
58126150	Turnover, meat- and cheese-filled, tomato-based sauce
58133130	Manicotti, cheese-filled, with meat sauce
58134610	Tortellini, meat-filled, with tomato sauce
58134710	Tortellini, spinach-filled, with tomato sauce

# Intertek

58140110	Spaghetti with corned beef, Puerto Rican style
58145140	Macaroni or noodles with cheese and tomato
58146300	Pasta, whole wheat, with meat sauce
58146310	Pasta, whole wheat, with tomato sauce, meatless
58155110	Rice with chicken, Puerto Rican style (Arroz con Pollo)
58155210	Stuffed rice with chicken, Dominican style (Arroz relleno Dominicano)
58155810	Stewed rice, Puerto Rican style (arroz quisado)
58155910	Rice with squid, Puerto Rican style (arroz con calamares)
58156210	Rice with vienna sausage, Puerto Rican style (arroz con salchichas)
58156310	Rice with Spanish sausage, Puerto Rican style
71931010	Cassava with creole sauce, Puerto Rican style (Yuca al mojo)
74415110	Tomato and sofrito stewing sauce, Puerto Rican style
75410500	Chiles rellenos, cheese-filled (stuffed chili peppers)
75412060	Eggplant parmesan casserole, regular
77141010	Potato chicken pie, Puerto Rican style (Pastelon de pollo)
77205110	Ripe plantain fritters, Puerto Rican style (Pionono)
77230510	Cassava pie stuffed with crab meat, Puerto Rican style (Empanada de jueyes)
77250110	Stuffed tannier fritters, Puerto Rican style (Alcapurrias)
77316010	Stuffed cabbage, with meat, Puerto Rican style (Repollo relleno con carne)
77316510	Stuffed cabbage, with meat and rice, Syrian dish, Puerto Rican style (Repollo relleno con carne y co)
77563010	Puerto Rican stew (Sancocho)

## Nut and Nut Products

### Nut butters and nut spreads

[EPG] = 0.21% (38% adjusted for fat portion of 55%)

42200500	Almond butter
42200600	Almond paste (Marzipan paste)
42201000	Cashew butter
42202000	Peanut butter
42202010	Peanut butter, low sodium
42202100	Peanut butter, reduced sodium and reduced sugar
42202130	Peanut butter, reduced sugar
42202150	Peanut butter, reduced fat
42202200	Peanut butter, vitamin and mineral fortified
42203000	Peanut butter and jelly
91304090	Topping, chocolate flavored hazelnut spread

### Mixed foods containing nut butters and nut spreads

*Adjusted for peanut butter content of ~17 to ~31%*

42301010	Peanut butter sandwich
42302010	Peanut butter and jelly sandwich
42303010	Peanut butter and banana sandwich

## Soft Candy

### Candy Bars

[EPG] = 0.04% (17% adjusted for fat portion of 24%)

91700500	M&M's Almond Chocolate Candies
91703070	Rolo



91703150	Toblerone, milk chocolate with honey and almond nougat
91703200	TWIX Caramel Cookie Bars (formerly TWIX Cookie Bars)
91703250	TWIX Chocolate Fudge Cookie Bars
91703300	TWIX Peanut Butter Cookie Bars
91703400	Whatchamacallit
91705010	Milk chocolate candy, plain
91705020	Milk chocolate candy, with cereal
91705030	Kit Kat
91705040	Chocolate, milk, with nuts, not almond or peanuts
91705050	Milk chocolate candy, with fruit and nuts
91705060	Milk chocolate candy, with almonds
91705070	Chocolate, milk, with peanuts
91705090	Chocolate candy with fondant and caramel
91705200	Chocolate, semi-sweet morsel
91705300	Chocolate, sweet or dark
91705310	Chocolate, sweet or dark, with almonds
91705400	Chocolate, white
91705410	Chocolate, white, with almonds
91705420	Chocolate, white, with cereal
91705430	Kit Kat White
91705500	Mexican chocolate (tablet)
91715100	SNICKERS Bar
91715200	Baby Ruth
91715300	100 GRAND Bar
91718100	Butterfinger
91718110	Butterfinger Crisp
91726130	MILKY WAY Bar
91726140	MILKY WAY MIDNIGHT Bar (formerly MILKY WAY DARK Bar)
91726150	MARS Almond Bar (formerly MARS bar)
91726420	3 MUSKETEERS Bar
91726425	3 Musketeers Truffle Crisp Bar
91731010	M&M's Peanut Chocolate Candies
91731060	M&M's Peanut Butter Chocolate Candies
91734100	Reese's Peanut Butter Cup
91734200	Reese's Pieces
91734300	Reese's Sticks
91734400	Reese's Fast Break
91734450	Reese's Crispy Crunchy Bar
91746100	M&M's Milk Chocolate Candies (formerly M&M's Plain Chocolate Candies)
91746200	M&M's Pretzel Chocolate Candies

**SUBMISSION END**