

Countess™ II FL Automated Cell Counter

For fluorescence and brightfield applications

Catalog Number AMQAF1000

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About this guide

Audience

This user guide is for laboratory staff operating, maintaining, and analyzing data using the Countess[™] II FL Automated Cell Counter.

Revision history

Revision	Date	Description
C.0	Aug 2015	Remove Countess™ II Cell Counter information, update procedures for the new SW release, update legal and regulatory information and branding
B.0	Dec 2014	Correct technical specification for cell size
A.0	Sep 2014	New user guide

User documentation The guides listed below are available for the Countess™ II FL Automated Cell Counter.

Guide	Pub. no.
Countess™ II FL Automated Cell Counters User Guide	MAN0010644
Countess™ II and Countess™ II FL Automated Cell Counters Quick Reference Card (QRC)	MAN0010826

Additional resources are available on the Countess $^{\scriptscriptstyle{\text{TM}}}$ Technical Resources page. Go to www.thermofisher.com/countess to access protocols, application notes, and tutorials.

Text and keyboard conventions

Text and keyboard conventions used in this user guide are listed below. For safety alert words and symbols used in this document, see page 4.

Convention	Use
Bold	Bold text indicates user action. For example:
	Press More.
•	Right arrow symbol (▶) indicates a menu choice, and separates successive commands you execute or select from a drop-down or shortcut menu. For example: Select More ▶ Adjust.

User attention words

Two user attention words appear this document. Each word implies a particular level of observation or action as described below.



Note: Provides information that may be of interest or help but is not critical to the use of the product.



IMPORTANT! Provides information that is necessary for proper instrument operation, accurate installation, or safe use of a chemical.

Safety alert words

Four safety alert words appear in this document at points where you need to be aware of relevant hazards. Each alert word—IMPORTANT, CAUTION, WARNING, DANGER—implies a particular level of observation or action, as defined below:



IMPORTANT! – Provides information that is necessary for proper instrument operation, accurate installation, or safe use of a chemical.



CAUTION! – Indicates a potentially hazardous situation that, if not avoided, may result in minor or moderate injury. It may also be used to alert against unsafe practices.



WARNING! – Indicates a potentially hazardous situation that, if not avoided, could result in death or serious injury.



DANGER! – Indicates an imminently hazardous situation that, if not avoided, will result in death or serious injury. This signal word is to be limited to the most extreme situations.

Except for **IMPORTANT!** safety alerts, each safety alert word in this document appears with an open triangle figure that contains a hazard symbol. These hazard symbols are identical to the hazard symbols that are affixed to the instruments (see "**Safety symbols**" in Appendix E).

1. Product information

Product contents

The Countess $^{\text{\tiny{TM}}}$ II FL Automated Cell Counter is shipped with the components listed below.

Component	Quantity
Countess™ II FL Automated Cell Counter (Cat. no. AMQAF1000)	1 each
Power Cord with 4 adaptor cords	1 each
(for U.S./Canada/Taiwan/Japan, Europe, or UK)	
Countess™ Cell Counting Chamber Slides (50 slides/box)	1 box
Countess™ II FL Disposable Slide Holder	1 each
Countess™ II FL Reusable Slide Holder	1 each
Countess™ II FL Light Cube Removal Tool	1 each
Countess™ II USB drive	1 each
Countess™ II FL Automated Cell Counter Quick Reference Card	1 each

Upon receiving the instrument

Examine the instrument carefully for damage incurred during transit. Ensure that all parts of the instrument, including accessories listed above, are included with the product. Damage claims must be filed with the carrier; the warranty does not cover in-transit damage.

See page 8 for instructions on installing the instrument.

Register your instrument

Visit www.thermofisher.com/registercountess to register your instrument. You will be asked to supply the serial number, your name, and your contact details. Registering your instrument ensures that you will receive notifications of software upgrades and information on new assays for use with the Countess™ II FL Automated Cell Counter.

Product description

Countess™ II FL Automated Cell Counter

The Counters^{$^{\text{M}}$} II FL Automated Cell Counter is a fully automated, 3-channel cell counter and assay platform that uses EVOS^{$^{\text{M}}$} light cube technology, state-of-the-art optics, and image analysis algorithms to analyze fluorescently labeled cells or trypan blue stained samples in suspension.

- The Countess[™] II FL Automated Cell Counter offers an intuitive user interface, and provides the option to save data and generate a report, which can then be transferred to a PC using the USB drive supplied with the instrument or available separately.
- The cells to be counted are loaded into the instrument either in disposable
 Countess™ Cell Counting Chamber Slides or in glass Countess™ II FL Reusable
 Slides (page 15). Each chamber slide contains two enclosed chambers to hold the
 sample to allow you to measure two different samples or perform replicates of
 the same sample.
- The instrument takes 10 seconds per sample for a typical cell count in the brightfield channel and is compatible with a wide variety of eukaryotic cells. In addition to cell count and viability, the Countess™ II FL Automated Cell Counter also provides information on cell size.
- In addition to the brightfield channel, the Countess™ II FL Automated Cell Counter can accommodate two interchangeable EVOS™ fluorescent light cubes (page 49), enabling it to be used for multiple-fluorescence research applications.
- When equipped with EVOS™ light cubes, the Countess™ II FL Automated Cell Counter can be used to perform fluorescence assays for cells in suspension, including simultaneous counts of cells stained with two different fluorescent dyes, GFP and RFP expression, apoptosis, and cell viability (live, dead, and total cells). These assays are compatible with a wide variety of eukaryotic cells.

Instrument exterior components





- 1 Touch-screen display: The 7-inch capacitive touch-screen display is the main user interface of the Countess™ II FL Automated Cell Counter. It contains the buttons for all instrument functions and displays data from the cell count.
- (2) (7) **USB ports:** The USB ports allow you to transfer and save the cell count data and image to an external computer for record keeping and printing purposes. You can use the USB drive supplied with the instrument or any other standard, FAT32-formatted USB drive for data transfer. If desired, you can plug in a USB mouse into the rear USB port for instrument control.
 - **Note:** The USB ports located in the front and the back of the instrument function the same. However, the first USB drive connected will be the preferred saving location and both USB drives cannot be accessed at the same time.
- 3 Slide port: The slide port is used to insert the analysis slide containing the sample into the counter. The Countess™ II FL instrument accepts both the disposable Countess™ Cell Counting Chamber Slides and the glass Countess™ II FL Reusable Slides via interchangeable, slide-specific carriers. For more information, see "Slide operation", page 17.
- **Back panel:** The back panel of the Countess™ II FL Automated Cell Counter allows access to the optional EVOS™ light cubes and provides storage for the light cube tool and the reusable slide carrier. The back panel is secured to the instrument by two captive ¼-turn fasteners.
- (5) **Power switch:** The ON/OFF rocker switch is the main power switch. It is not necessary to use the power switch for day-to-day operation of the instrument.
- 6 **EVOS™ light cubes:** The EVOS™ light cubes allow the Countess™ II FL Automated Cell Counter to analyze fluorescently labeled samples. The Countess™ II FL Automated Cell Counter can accommodate two fluorescent light cubes. For more information, see "EVOS™ light cubes", page 49.
- 8 **Power input jack**: The power input jack connects the instrument to an electrical outlet through the supplied power cord and the appropriate plug, based on the electrical outlet configuration in your country.

2. Getting started

Installation

Operating environment

- Place the instrument on a level surface away from vibrations emanating from other pieces of equipment.
- Allow at least 5 cm (2 in) free space at the back of the instrument to allow for proper ventilation and prevent overheating of electronic components.
- Set up the instrument away from direct light sources, such as windows.

 Ambient room lighting can enter the imaging path and affect the image quality.
- Operating temperature range: 4°–32°C (40°–90°F).
- Relative humidity range: 30–90%.



IMPORTANT! Do not position the instrument so that it is difficult to turn off the main power switch located on the back of the instrument (see page 7). In case of an instrument malfunction, turn the main power switch to the OFF position and disconnect the instrument from the wall outlet.

Install the instrument

- 1. Unpack the instrument and place the instrument on a flat, level, dry surface.
- 2. Remove the thin plastic protector film from the touch-screen display.
- 3. Plug one end of the power cord appropriate for your region into the instrument.
- 4. Plug the power cord into the electrical outlet. Be sure to use only the power cord supplied with your instrument. Powering the instrument with an unapproved power cord may damage the instrument.

Turn ON the instrument

1. Turn on the instrument by flipping the **power switch** on the back of the instrument (page 7) to the **ON** position.

The instrument initializes and displays the Home screen.



- **2.** From the Home screen, you can proceed immediately to the assays by inserting a slide (page 19).
 - Alternatively, you can change or add a profile (Step 3, below) or change instrument settings (Step 4, below).
- 3. To change the current profile or to add a new profile to the instrument, press the **Profiles** button in the upper left corner.



- Profiles allow you to create customized count preferences (i.e., gate counts based on cell size, brightness, circularity, and/or relative fluorescence intensity) (see page 10).
- **4.** To change instruments settings, press the **Instrument Settings** button in the upper right corner.

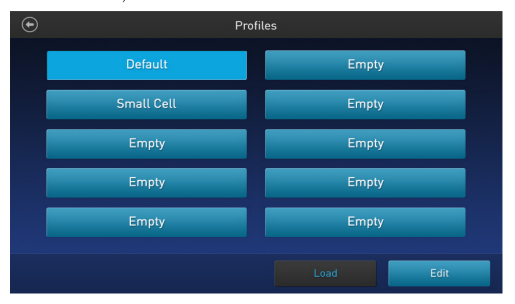


Instrument settings allow you to update the CountessTM II software, change the date and time, and install or change up to two EVOSTM light cubes (page 36).

Load profile

Profiles screen

Profiles screen allows you to create and save up to 9 customized profiles. Each custom profile defines the count parameters (size, brightness, circularity, and fluorescence intensity) and automatic instrument functions (Auto Lighting and Auto FL Threshold) for a consistent and streamlined workflow.



- You can access the Profiles screen from the Home, Capture, Results, Advanced, or Adjust screens.
- The current profile is displayed on the upper left corner of the Home, Capture, Results, Advanced, or Adjust screens.



- Automatic instrument functions (below) and count parameters (page 11) are defined in the Edit profile screen (see "Add/edit a profile", page 12).
- The Default profile contains default count settings and cannot be edited.
- The count parameters specified in the selected profile are applied to all new cell counts.
- If you have already performed a count, loading a new profile from the Results screen applies the count preferences to the current counts results (total cells, viability etc.) and to all new counts.

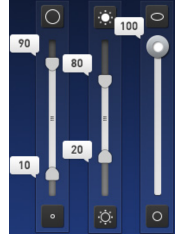
Automatic instrument functions

Auto FL Threshold and Auto Lighting functions are turned on and off in the Edit profile screen using the **Auto FL Threshold** and **Auto Lighting** checkboxes (see page 12).

- **Auto FL Threshold:** Automatically applies threshold in fluorescence channels to subtract background fluorescence for improved analysis despite variable background levels between samples. This function is available only for instruments equipped with the optional EVOS™ light cubes.
- **Auto Lighting:** Automatically illuminates the sample in the brightfield for increased sample-to-sample consistency and decreased user-to-user variability.

Count parameters

Count parameters are adjusted in the Edit profile screen using the **parameter sliders**. Parameter sliders correspond to a single channel, which is selected using the **channel selection** radio buttons located above the sliders.





Size, brightness, and circularity sliders

Fluorescence intensity slider

- **Size:** As you move the slider up, the algorithm includes larger objects in the count. As you move the slider down, only the smaller objects are counted.
 - = larger objects
- = smaller objects
- **Brightness:** As you move the slider up, the algorithm includes the brighter objects in the count. As you move the slider down, only the dimmest of objects are counted.
 - = brighter objects
- = dimmer objects
- Circularity: As you move the slider up, the algorithm includes more objects
 with shapes other than circular in the count. As you move the slider down,
 only the objects that are perfect circles are counted.
 - = less circular
- o = more circular
- Fluorescence intensity: As you move the slider up, the algorithm includes the
 objects that fluoresce more brightly in the count. As you move the slider down,
 only the dimmest of objects are counted.
 - = brightly fluorescent objects = dim or less fluorescent objects
- Size, brightness, and fluorescence intensity sliders are range sliders.
 - To adjust the upper and lower boundaries without changing the data range, drag the slider by its middle section (i.e., the slider bar).
 - To adjust only the upper or the lower boundary, move the upper or the lower handle in the desired direction. This will also change the range of values within which the cells are counted.
- The **circularity** slider only sets a single threshold value; cells that fall below the set value are counted, and cells that are beyond this range are excluded.
 - To adjust the threshold for circularity, drag the slider in the desired direction.

Load a profile

1. Press the **Profiles** button located on the upper left corner of the screen to open the Profiles screen.



- **2.** Press the desired profile to select, and then press **Load**.
 - The instrument will load the count parameters specified in the selected profile and return to the previous screen.
- **3.** To return to the previous screen without loading the new profile, press the **previous** button.



The instrument will keep the saved profile, but return to the previous screen without loading it.

Add/edit a profile

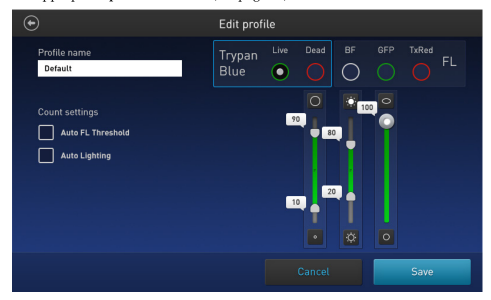
1. Press the **Profiles** button located on the upper left corner of the screen to navigate to the Profiles screen.



- 2. To add or edit a new profile, select an empty or an existing profile, and then press **Edit**. The Edit screen for the selected profile opens.
 - **Note:** The Default profile contains default count settings and cannot be edited.
- **3.** Select or deselect the **Auto FL Threshold** checkbox to turn the Auto FL Threshold function **ON** or **OFF** (page 10).
 - **Note:** This function is available only for instruments equipped with the optional EVOSTM light cubes.
- **4.** Select or deselect the **Auto Lighting** checkbox to turn the Auto Lighting function **ON** or **OFF** (page 10).

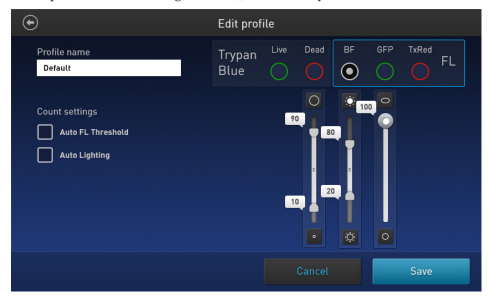
Define count parameters for the brightfield channel:

- **5**. To define the new count parameters in the brightfield channel:
 - **a.** From the **Trypan Blue** selection box, select **Live** or **Dead** radio buttons (see image below).
 - **b.** Adjust the size, brightness, and circularity thresholds using the appropriate **parameter slider** (see page 11).



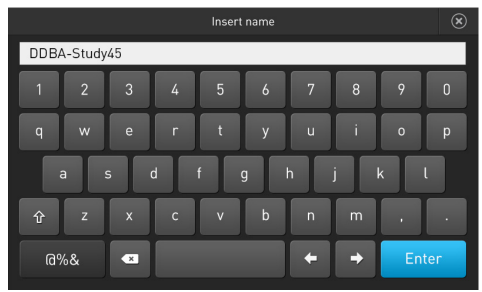
Define count parameters for the fluorescent channels

- **6.** To define the new count parameters for fluorescence assays (available only for instruments equipped with the optional EVOS™ light cubes):
 - **a.** From the **FL** selection box, select the desired channel using the appropriate radio button (see image below).
 - The available options are **BF** (brightfield) and up to two fluorescence channels, depending on the light cube(s) installed (**GFP** and **TxRed** in the example below).
 - **b.** Define the new count parameters for the selected channel using the appropriate **parameter slider** (see page 11).
 - **c.** Repeat for the remaining channel(s), as necessary.



Note: Parameter sliders in the BF channel allow you to gate count results based on size, brightness, and circularity. The parameter slider in the selected fluorescence channel allows you to gate count results based on relative fluorescence intensity in that channel.

7. To assign a name to the new profile or to change the name of the existing profile, press the **Profile name** text box. The alpha-numeric **keypad** opens.



- **8.** Type in the desired profile name using the alpha-numeric keypad. To enter symbols, press the **symbol** (@%&) key. To return to the alphanumeric keypad, press **ABC**.
- 9. Press Enter to save the name and return to the Edit profile screen.
 To return to the Edit profile screen without saving the name, press the close button.



- **10.** Press **Save** to save the new profile, and then press **Close** in the confirmation screen to return to the Profiles screen.
 - To return to the Profiles screen without saving, press Cancel.
- 11. On the Profiles screen, press **Load**. The instrument will load the count parameters specified in the selected profile and return to the previous screen.
- **12**. To return to the previous screen without loading the new profile, press the **previous** button. The instrument will keep the saved profile, but return to the previous screen without loading it.



Prepare sample

Recommendations

To obtain the best results, follow these recommendations:

- Ensure that the cell sample is homogeneously mixed.
- The measurement range extends from 1×10^4 – 1×10^7 cells/mL, but the optimal range is 1×10^5 – 4×10^6 cells/mL.
- For accurate results in cell viability assays, ensure that the counting area is covered with the cell suspension and count the cells immediately after staining per the assay protocol.
- Do **not** press the optical surfaces of the chamber slides. Hold the slides by the edges.
- Take care to avoid forming bubbles in the sample.

Load Countess™ Chamber Slide

- 1. Prepare the sample by adding 10 μ L of your cell suspension to 10 μ L of 0.4% trypan blue stain. Mix the sample mixture well by pipetting it up and down a few times.
- 2. Gently pipet 10 μ L of the sample into the half moon-shaped sample loading area. The sample is loaded into the chamber through capillary action.



- **3.** Let the sample mixture settle in the chamber for 30 seconds, and then insert the slide into the slide port (see page 7). You will hear a soft click, if the slide is pushed in correctly.
- 4. To remove the slide, push the slide gently into the instrument until it "clicks" and a spring pushes the slide out. Grasp the slide and pull it out the rest of the way.



Note: After using the Countess[™] Cell Counting Chamber Slides, appropriately dispose of them as biohazardous waste. Do **not** reuse the disposable chamber slides.

Load Countess™ II FL Reusable Slide

- Before loading your sample into the Countess™ II FL Reusable Slide, place a
 cover slip on the counting chamber, making sure the cover slip is clean and
 free of grease.
- 2. Gently pipet $10 \mu L$ of the sample into the sample inlet, allowing capillary action to draw the sample into the counting chamber. A properly loaded counting chamber should have a thin, even film of fluid under the cover slip.



3. After using the Countess™ II FL Reusable Slide, rinse the glass slide and cover slip with water, and then clean with 70% ethanol. Use Kimwipes™ laboratory tissues to clean and dry the slides, as needed.



Note: Each chamber in the CountessTM Cell Counting Chamber Slide or the CountessTM II FL Reusable Slide has a $10-\mu$ L sample capacity. Do not overfill the slide chambers.

Slide operation

The Countess^{$^{\text{TM}}$} II FL instrument accepts both disposable Countess^{$^{\text{TM}}$} Cell Counting Chamber Slides and glass Countess^{$^{\text{TM}}$} II FL Reusable Slides on interchangeable, slide-specific carriers.

Countess™ Cell Counting Chamber Slide

1. To use the plastic, disposable Countess[™] Cell Counting Chamber Slide with the Countess[™] II FL Automated Cell Counter, insert the slide carrier (black, see image below) into the slide port of the instrument until it clicks into place.

Note: The Countess[™] II FL Automated Cell Counter is shipped with the disposable slide carrier already installed



2. Load the chamber slide with your sample as described on page 15, and then insert the slide into the slide carrier in the slide port until it clicks into place.



- **3.** To remove the slide, push the slide gently into the instrument until it "clicks" and a spring pushes the slide out. Grasp the slide and pull it out the rest of the way.
- **4.** *Optional*: To remove the slide carrier, gently squeeze the tabs and pull the carrier completely out of the instrument.

Note: You can store the slide carrier behind the access panel on the back of the instrument (see page 7).

Countess™ II FL Reusable Slide

1. To use the Countess™ II FL Reusable Slide, unlatch the back panel of the Countess™ II FL Automated Cell Counter with the two captive ¼-turn fasteners that secure the back panel on the rear of the instrument.



2. Remove the reusable slide carrier (white) from inside of the back panel.



3. Load the reusable glass slide with the sample as described on page 16, and place the loaded slide into the white slide carrier.



- **4.** Insert the carrier and reusable slide assembly into the slide port, and gently push into the instrument until it "clicks" into place.
- **5.** To remove the slide, push the slide gently into the instrument until it "clicks" and a spring pushes the slide out. Grasp the slide and pull it out the rest of the way.
- **6.** *Optional*: To count the second sample present on the reusable slide, simply remove the slide from the carrier, rotate, and reinsert the slide into the carrier so that the second sample is aligned with the sample viewing hole.

Note: You can store the slide carrier behind the access panel on the back of the instrument (see page 7).

3. Cell count and cell viability assays

Count cells in brightfield

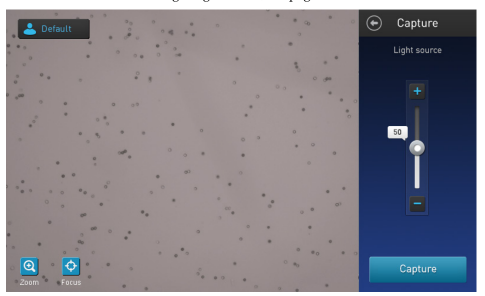
Count procedure

- 1. Prepare the sample by adding $10~\mu L$ of your cell suspension to $10~\mu L$ of 0.4% trypan blue stain. Mix the sample mixture well by pipetting up and down a few times.
- 2. Load 10 μ L of the sample mixture per chamber into the sample slide as described on page 15. Let the sample mixture settle for 30 seconds.
- **3.** *Optional*: Press the **Profiles** button and load the desired profile as described on page 12.



- **4.** Insert the sample slide into the slide port (see page 7), making sure that the sample side is inserted completely into the instrument. You will hear a soft click, if the slide is pushed in correctly.
- **5**. When the slide is inserted, the instrument automatically illuminates the sample, sets the intensity of brightfield illumination, and auto focuses on the cells.

Note: To turn off the Auto Lighting function, see page 13.



6. *Optional*: To manually adjust the focus, press the **Focus** button, and then use the **Focus slider** to bring your sample into focus as described on page 44.



7. Press the **Set** button to set the focus and collapse the focus controls. Once the focus has been set, the Set button on the focus slider becomes inactive, confirming that the focus setting has been stored.



Note: If needed, **Zoom** in on the image to adjust focus or lighting.

⊕

8. *Optional*: Set exposure using the **light source slider**.

The light source slider controls the LED intensity, camera gain, and exposure time and it is used for adjusting the image brightness.

Note: If your instrument is equipped with an EVOS[™] light cube, first press **Adjust**, and then select **brightfield** (white circle) as the light source. Set the exposure, then press **Done** to return to the Capture screen.

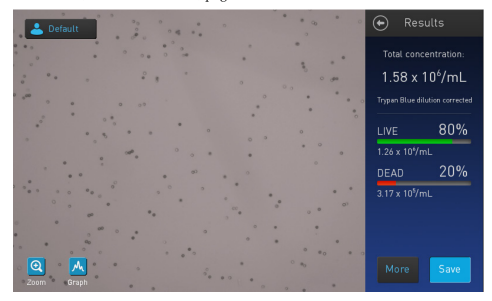


9. Press Capture.

Note: If your instrument is equipped with an EVOS[™] light cube, make sure that only the **BF** (brightfield) checkbox is selected under Collect channels before capturing the image.



The instrument temporarily captures the image and displays the results (total concentration, percentage and concentration of live and dead cells). For more information, see "View results", page 21.



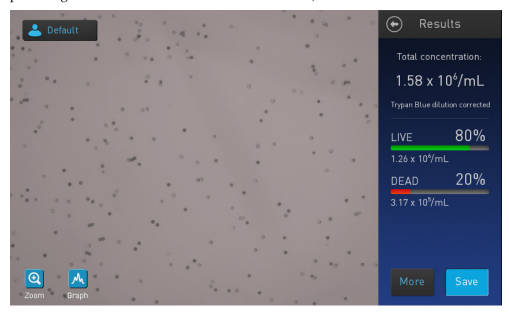
Next steps

- To identify the objects (i.e., cells) counted as "Live" or "Dead", press **More** to navigate to the Advanced screen (page 22).
- To see the distribution of live and dead cells in a graphical format, press the **Graph** button (see page 23).
- To gate the results by object size, brightness, or circularity, first press **More** to open the Advanced screen, and then press **Adjust** to navigate to the Adjust screen (page 24).
- To permanently save the results, press **Save** (see page 33).
- To perform a new count, remove the slide and reinsert it second chamber first into the instrument, or insert a new sample slide.

View results

Results screen for brightfield

The Results screen for cell count and cell viability assays performed using the brightfield channel displays a composite image of the objects counted and the results of the cell count and cell viability calculations (total concentration, percentage and concentration of live and dead cells).





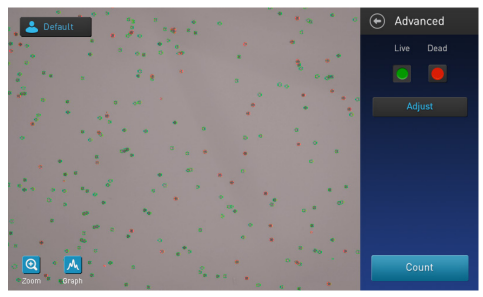
Note: When performing cell counts and cell viability assays in brightfield, the counting algorithm assumes that you have diluted your cells 1:1 in trypan blue and takes this dilution into account when calculating the total cell concentration. The cell concentration displayed in the Results screen is the original cell concentration before dilution into trypan blue.

Identify objects counted

Advanced screen

The Advanced screen allows you to identify the objects (i.e., cells) counted in each channel and included in the count results for further review. After reviewing the marked objects, you can adjust the threshold for size, brightness, and/or circularity as desired for your application.

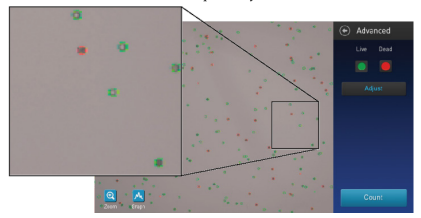
Identify cells counted in cell count and cell viability assays 1. On the Results screen, click **More.** The Advanced screen opens.



2. To identify the cells that are included in the count as "live", press the **Live** button. "Live" cells will be circled in green on the screen.

To identify the cells that are included in the count as "dead", press the **Dead** button. "Dead" cells will be circled in red on the screen.

Note: You may select either or both options. In the example below, both **Live** and **Dead** buttons are pressed and "live" and "dead" cells are marked with green and red circles around them, respectively.



3. To unmark the cells identified as "live" (green) or "dead" (red) on the screen, press the **Live** or the **Dead** button again, respectively.

Graph count results

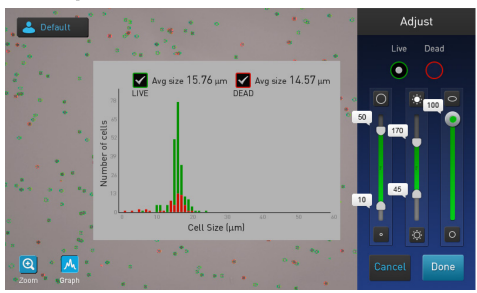
View graph

For cell count and cell viability assays performed in the brightfield channel, you can view the distribution of cells (live and/or dead) based on size in a graphical format.

Note: You can view the Graph on Results, Advanced, and Adjust screens.

1. To view the graph showing the distribution live and/or dead cells based on cell size, press the **Graph** button.





- 2. To view the distribution of only the live or dead cells, check the corresponding **Live** or **Dead** check box on the graph.
 - The graph will automatically update and display the distribution of cells based on size only in the selected population.
- **3.** *Optional*: Using the **size**, **brightness**, and **circularity** sliders, adjust the count parameters. As you adjust the count parameters, the count results and the graph will be automatically updated.
- **4.** To close the graph, press the **Graph** button again.

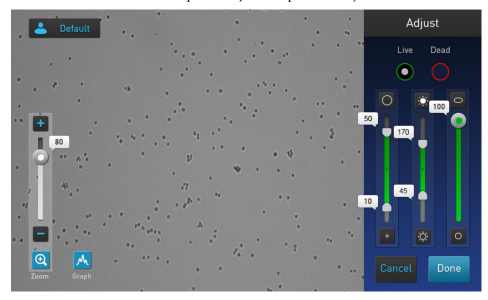
Gate count results

Adjust screen

The Adjust screen for cell count and cell viability assays in the brightfield channel contains the controls for gating count results based on size, brightness, and circularity. You can adjust the count parameters before or after performing a count.

Gate count results

- 1. On the Results screen, press **More** to open the Advanced screen.
- **2.** *Optional*: Press the **Live** and/or the **Dead** button to identify the cells in the selected population (see page 22).
- 3. On the Advanced screen, press **Adjust** to open the Adjust screen.



4. *Optional*: Press the **Graph** button to view the distribution of cells (live and/or dead) based on size as you gate the count results (see page 24).



- 5. Select the channel (Live or Dead) you wish to gate.
- **6.** Using the **size**, **brightness**, and **circularity sliders**, adjust the count parameters. **Note**: For a description of the count parameters and count parameter controls (i.e., parameter sliders), see page 11.
- 7. When finished, press **Done** to save the changes to count parameters and return to the Advanced screen.
 - Press Cancel to return to the Results screen without saving the changes.
- **8.** On the Advanced screen, press **Count** to recalculate your results with the new count parameters.
- **9.** To permanently save your results, see page 33.

4. Fluorescence assays

Count cell fluorescence

Overview

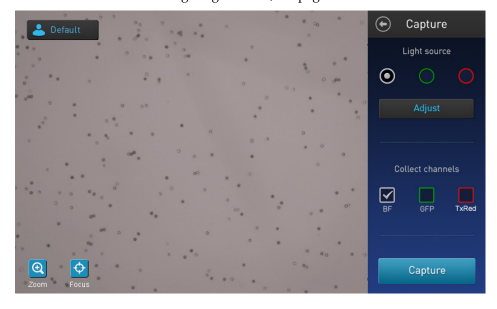
Countess™ II FL Automated Cell Counter equipped with the optional EVOS™ light cubes can be used for a variety of fluorescent applications, including simultaneous counts of cells stained with two different fluorescent dyes, GFP and RFP expression, and apoptosis and cell viability assays.

For instructions on installing EVOSTM light cubes to your CountessTM II FL Cell Counter, see page 40.

Count procedure

- 1. Ensure that your fluorescent cell sample is homogeneously mixed.
- 2. Load $10 \mu L$ of the fluorescent sample mixture per chamber into the sample slide as described on page 15. Let the sample mixture settle for 30 seconds.
- **3.** *Optional*: Press the **Profiles** button located on the upper left corner of the screen to open the Profiles screen and load the desired profile as described on page 10.
- **4.** Insert the sample slide into the slide port (see page 7), making sure that the sample side is inserted completely into the instrument. You will hear a soft click, if the slide is pushed in correctly.
- **5**. When the slide is inserted, the instrument automatically illuminates the sample, sets the intensity of brightfield illumination, and auto focuses on the cells.

Note: To turn off the Auto Lighting function, see page 13.

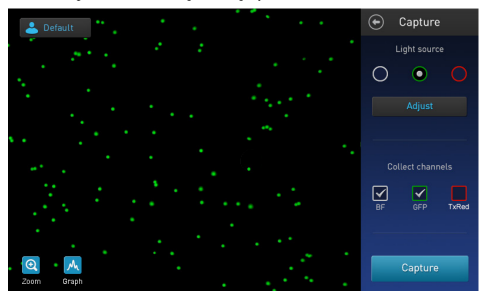


- **6.** *Optional*: To manually adjust the focus, press the **Focus** button and use the **Focus slider** to bring your sample into focus as described on page 44.
- ф
- 7. Press the **Set** button to set the focus and collapse the focus controls. Once the focus has been set, the Set button on the focus slider becomes inactive, confirming that the focus setting has been stored.



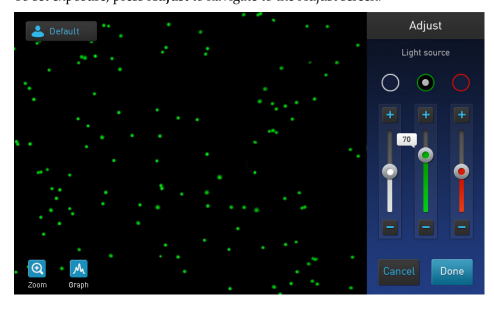
8. To view your sample under a different light source, press the desired **Light source** button. The instrument displays the sample in the selected channel (brightfield or fluorescent).

In the example below, the sample is displayed in the GFP channel.



Note: The light source buttons select the light channel (brightfield and/or fluorescence) for sample illumination and are used when setting the exposure for the selected channel (see Steps 9–11); they do not determine which channels are used for capturing the image.

9. To set exposure, press **Adjust** to navigate to the Adjust screen.

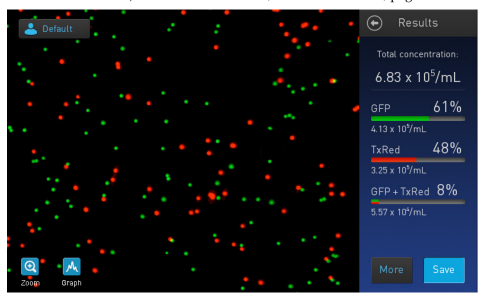


- **10.** Press the **light source** button for the channel you wish to set exposure and adjust the exposure using the **light source slider**. Repeat the procedure for the remaining channels, if desired.
- **11.** After setting the exposure, press **Done** to return to the Capture screen. To return to the Capture screen without changing the exposure, press **Cancel**.
- **12.** On the Capture screen, select the **Collect channels** check boxes for the channels you wish to capture.



13. Press Capture.

The instrument temporarily captures the image and displays the results (total concentration, percentage and concentration of cells counted in each fluorescence channel). For more information, see "View results", page 28.



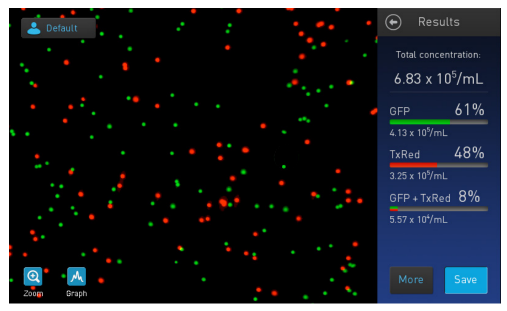
Next steps

- To identify the objects (i.e., cells) counted in each channel, press **More** to navigate to the Advanced screen (page 29).
- To see the distribution of cells counted through each channel in a graphical format, press the **Graph** button (see page 30).
- To gate the results by object size, brightness, circularity, or relative fluorescence intensity, first press **More** to open the Advanced screen, and then press **Adjust** to navigate to the Adjust screen (page 32).
- To permanently save the results, press **Save** (see page 33).
- To perform a new count, remove the slide and reinsert it second chamber first into the instrument, or insert a new sample slide.

View results

Results screen for cell fluorescence assays

The Results screen for cell fluorescence assays displays a composite image of the objects counted and the results of the cell count and cell viability calculations (total concentration, percentage and concentration of cells counted through each fluorescence channel).





Note: The total cell concentration displayed after a fluorescent count does not take any dilution into account. Therefore, the results reflect the actual cell concentration in the sample slide, which must be multiplied by any dilution factor present to calculate the original cell concentration.

This is in contrast to the cell counts in brightfield, where the counting algorithm assumes a 1:1 dilution of the sample in trypan blue and displays the original cell concentration (i.e., before the dilution) in the Results screen.

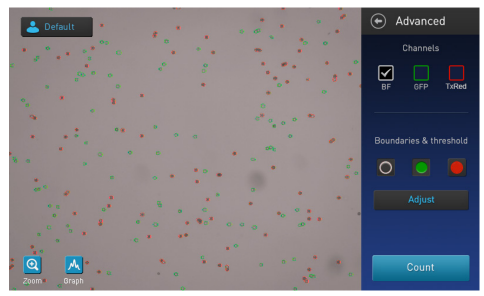
Identify objects counted

Advanced screen

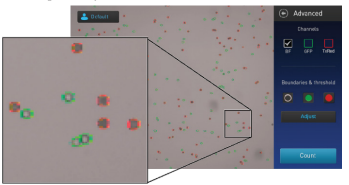
The Advanced screen allows you to identify the objects (i.e., cells) counted in each channel and included in the count results for further review. After reviewing the marked objects, you can adjust the threshold for size, brightness, and/or circularity as desired for your application.

Identify cells counted in fluorescence assays

1. On the Results screen, click **More** to open the Advanced screen.



- **2.** *Optional*: To view your sample under a specific light source (brightfield and/or fluorescent), select the desired **Channels** checkbox (brightfield in the example above). You may display your sample in any or all of the available channels.
- **3.** To identify the cells that are counted in a specific channel, press the corresponding **boundaries** button. Cells counted in the selected channel will be circled on the screen with the same color as the selected channel.
 - In the example below, both the **GFP** and **TxRed boundaries** buttons are pressed and the cells counted in the GFP and TxRed channels are marked with green and red circles, respectively.



4. To unmark the cells counted in a specific channel, press the corresponding **boundaries** button again.

Graph count results

View graph for cell fluorescence assays

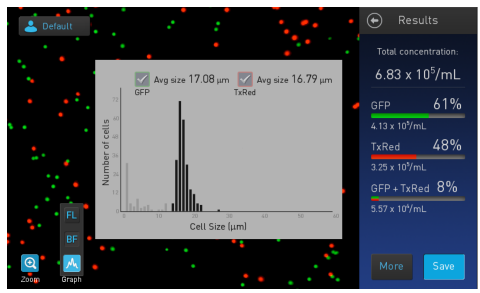
For fluorescence assays, you have the option of viewing the distribution of the cells based on size or based on relative fluorescence intensity in a graphical format.

Note: You can view the Graph on Results, Advanced, and Adjust screens.

1. To view the graph showing the distribution of cells based on size, press the **Graph** button, and then select **BF** (brightfield).

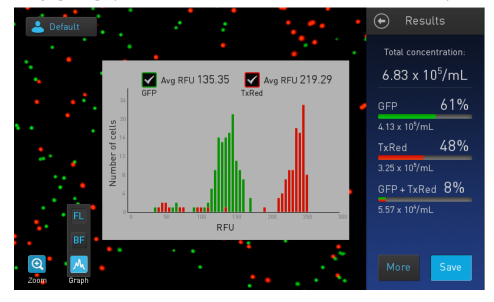
The graph displays the size distribution of the total cell count (number of cells vs. cell size in μm), and the average size of the cells counted in each available fluorescence channel.





2. To view the distribution of cells based on relative fluorescence intensity, press the **Graph** button, and then select **FL** (fluorescence).

The graph displays the distribution of cells based on fluorescence intensity.



- **3.** To remove the cells counted in a specific channel from the graph, uncheck the corresponding **channel** check box on the graph.
 - The graph automatically updates and displays the distribution of cells based on relative fluorescence intensity only in the selected (i.e., checked) channel.
- **4.** To add the cells counted in a specific channel to the graph, re-check the corresponding **channel** check box.
- **5**. To close the graph, press the **Graph** button again.

Gate count results

Adjust screen

The Adjust screen for cell fluorescence assays contains the controls for gating count results based on size, brightness, circularity, and fluorescence intensity. You can adjust the count parameters after performing a count.

Gate count results

- 1. On the Results screen, press **More** to open the Advanced screen.
- **2.** *Optional*: To view your sample under a specific light source (brightfield and/or fluorescent), select the desired **Channels** checkbox on the Advanced screen (see page 29).
- **3.** *Optional*: Press the desired **boundaries** button(s) on the Advanced screen to identify the cells counted in the corresponding channel (see page 29).
- **4.** Press **Adjust** to open the Adjust count parameters screen, which contains the controls for adjusting the count parameter(s) in the selected channel.

Note: For a description of the count parameters and count parameter controls (i.e., parameter sliders), see page 11.



5. *Optional*: Press the **Graph** button to view the distribution of cells based on size or fluorescence intensity (see page 30).



- **6.** Select the **brightfield channel** (white circle) to adjust the thresholds for size, brightness, and circularity using the **size**, **brightness**, and **circularity** sliders.
- **7.** Select the desired **fluorescence channel** (colored circles) to adjust the threshold for fluorescence intensity using the **fluorescence intensity** slider.
 - **Note:** The fluorescence channels available depend on the EVOS[™] light cubes installed in the instrument.
- **8.** When finished, press **Done** to save the changes to count parameters and return to the Advanced screen.
 - Press Cancel to return to the Results screen without saving the changes.
- **9.** On the Advanced screen, press **Count** to recalculate your results with the new count parameters.

6. Save results

Save count results

Save screen

The Countess™ II FL Automated Cell Counter allows you to save your data and images using a USB flash drive.

To save your experiment, you can choose from the following options, in any combination:

- **Result:** Saves the Results screen as it is displayed on the instrument, with or without the Graph, in the selected image format (JPEG, BMP, PNG, or TIFF).
- **Images:** Saves only the raw captured image in the selected image format (JPEG, BMP, PNG, or TIFF).
- Data: Saves the data from the experiment as a CSV file (comma separated values). The CSV format allows for processing or re-displaying results with any third party software or spreadsheet program. For more information on the CSV file format, see "Appendix D: CSV file format", page 51.



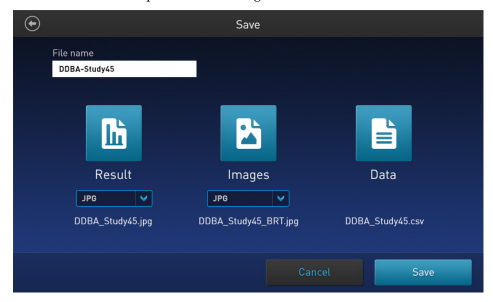
Note: If you wish to save your results with the Graph showing the distribution of cells based on cell size or fluorescence intensity, make sure that the desired graph is displayed on the Results screen.

Save procedure

1. To save your data, insert the Countess[™] II USB drive (or equivalent) into an available USB port on the instrument (see page 7).

Note: The USB ports located in the front and the back of the instrument function the same. However, the first USB drive connected will be the preferred saving location.

2. On the Results screen, press **Save** to navigate to the Save screen.

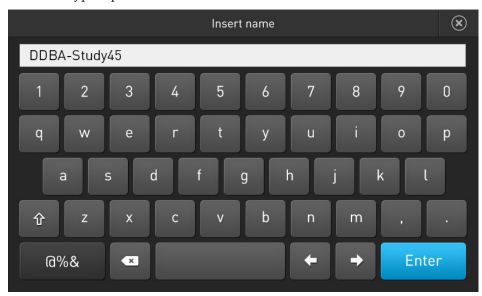


3. To assign a name to your experiment, press the **File name** text field. The alpha-numeric **keypad** opens.



4. Enter the file name using the alpha-numeric **keypad**.

To enter symbols, press the **symbol** (@%&) key. To return to the alphanumeric keypad, press **ABC**.

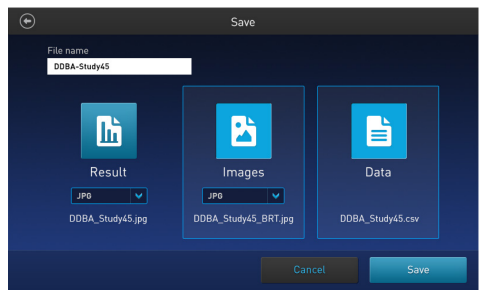


Press Enter to save the name and return to the Save screen.To return to the Save screen without saving the name, press the close button



6. Select the desired mode(s) to save your experiment (**Result, Images, Data**). You can select an individual mode (e.g., Result only) or any combination of modes (e.g., Result, Images, and Data).

In the example below, **Images** and **Data** are selected.



By default, Result and Images are saved as JPEG files.
 To choose a different file format, press the file type button.
 Choose file type screen opens.



Note: Data can only be saved as a CSV file.

8. Press to select the desired **file type**. Available options are **JPEG**, **BMP**, **PNG**, and **TIFF**.

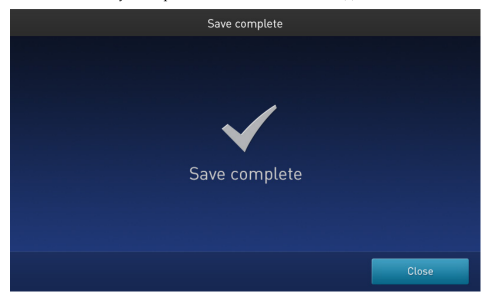


After you make your selection, the instrument returns to the Save screen.

To return to the Save screen without changing the file format, press the **close** button.



9. Press **Save** to save your experiment in the selected mode(s) in the USB drive.



10. Press Close and then transfer the USB drive to the desired location.

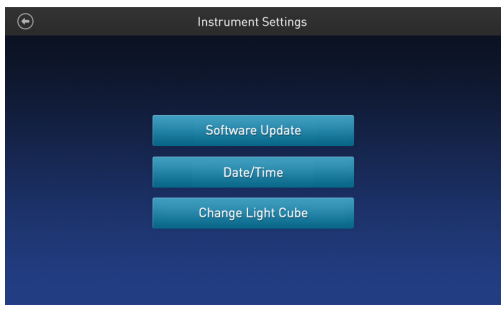
7. Instrument settings

Overview

Instrument Settings screen

To access the Instrument Settings screen, press the **Instrument Settings** button on the Home page (page 9).





In the Instrument Settings screen, you can:

- perform software update (page 37)
- set the date and time (page 38)
- change or install EVOS[™] light cube (page 40)

Software update

Guidelines for software update

• The USB drive used for transferring the software update file must be FAT32 formatted; verify this before proceeding. If necessary, reformat the USB drive to FAT32 following the recommended procedure for your operating system.

Note: Reformatting the USB drive will result in the loss of all files. Back up the files in the USB drive prior to reformatting.

- The software update file must be saved on the top level of the USB drive, not within a folder or a subfolder.
- The software update file must be uncorrupted during transfer. Do not rename, zip, or compress the software update file.

Update the Countess™ II/II FL software

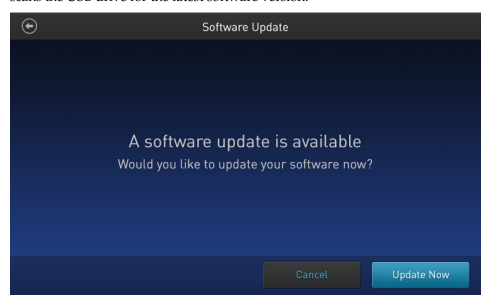
1. Go to **www.thermofisher.com/countessupdate**, and download the latest Countess[™] II/II FL software version to your desktop.

Note: The software update file has a version-specific name followed by the extension .lft (e.g., Countess_II_v_1_0_202.lft for software version 1.0.202).

- **2.** Copy the software update file onto the USB drive, making sure that it is saved on the top level and not hidden within a folder.
- 3. Insert the USB drive into one of the USB ports of the instrument (see page 7).
- **4.** Press the **Instrument Settings** button on the Home page (page 9) to open the Instrument Settings screen (page 36).



5. Select **Software Update** from the Instrument Settings menu. The instrument scans the USB drive for the latest software version.



- When prompted, select Update Now.
- 7. Once the update has completed, restart the instrument.

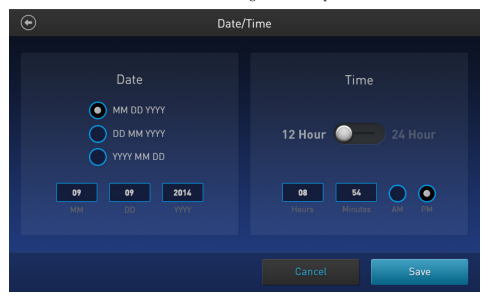
Date/Time

Set the date and time

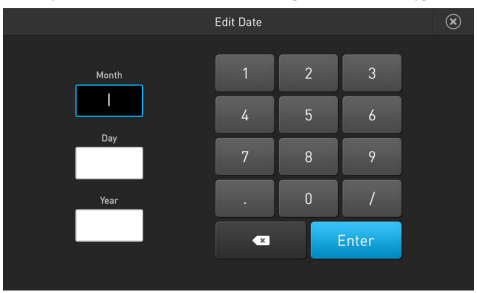
1. Press the **Instrument Settings** button on the Home page (page 13) to open the Instrument Settings screen.



2. Press **Date/Time** on the Instrument Settings menu to open the Date/Time screen.



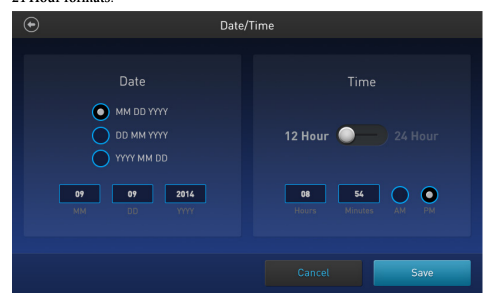
- 3. Select the **Date format** you wish to use.
- 4. Press any **Date** text box (**MM**, **DD**, or **YYYY**) to open the **Edit Date** keypad.



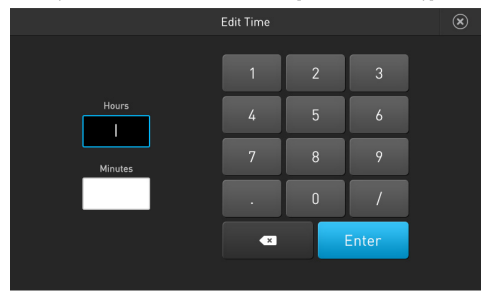
- 5. Using the keypad, enter the date into **Month**, **Day**, and **Year** text boxes, pressing **Enter** after each entry.
- **6.** After you are finished entering the date, press the **close** button to return to the Date/Time screen.



7. Select the **Time format** you wish to use. Available options are **12 Hour** and **24 Hour formats**.



8. Press any Time text box (Hours or Minutes) to open the Edit Time keypad.



- **9.** Using the keypad, enter the time into **Hours** and **Minutes** text boxes, pressing **Enter** after each entry.
- **10.** After you are finished entering the time, press the **close** button to return to the Date/Time screen.



- 11. If you have selected the 12 Hour format, select AM or PM.
- **12.** Press **Save** to set the Time and Date and return to the Instrument Settings

Press **Cancel** to return to the Instrument Settings screen without saving your changes.

Change light cube

Install or change EVOS™ light cube The Countess^{$^{\text{M}}$} II FL Automated Cell Counter can accommodate up to two EVOS^{$^{\text{M}}$} light cubes. Each user-interchangable, auto-configured EVOS^{$^{\text{M}}$} light cube contains an LED, collimating optics, and filters for fluorescence applications. EVOS^{$^{\text{M}}$} light cubes do not come standard with the device and must be purchased separately (see page 49). To install or change a light cube:

1. Press the **Instrument Settings button** on the Home page (page 9) to open the Instrument Settings screen.



- 2. Press **Change Light Cube**. The instrument positions the light cube tray to enable light cube installation.
- **3.** When prompted, power off the Countess[™] II FL Automated Cell Counter using the **power switch** on the back of the instrument (page 7).
- **4.** Unplug the power cord from the Countess[™] II FL Automated Cell Counter.
- 5. Unlatch the back panel with the two captive ¼-turn fasteners (indicated by red arrows) that secure the back panel on the rear of the Countess™ II FL Automated Cell Counter and remove the back panel.



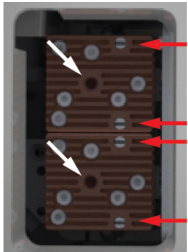
6. Place the light cube into one of the empty slots in the back of the device.



7. Using the tool provided on the inside of the back panel (Figure A, below), secure the light cube by tightening the two screws on the end of the cube (red arrows in Figure B, below).

A B





- **8**. To remove a light cube, unscrew both screws that secure it to the instrument.
- Thread the light cube removal tool into the central hole in the cube (white arrows in Figure B, above) and gently pull the light cube out of the device.
 Note: Always store the cube removal tool in the back panel for easy access.
- 10. Install the back panel and secure it in its place with both ¼-turn fasteners.
- 11. Plug the power cord back into the Countess[™] II FL Automated Cell Counter.
- **12.** Turn off the Countess[™] II FL Automated Cell Counter by flipping the **power switch** on the back of the instrument to the ON position.

8. Maintenance

Instrument care

General guidelines for care

- Use the appropriate cleaning solutions for each component, as indicated in the cleaning procedures on page 43.
- If liquid spills on the instrument, turn off the power immediately and wipe dry.

Power supply

Always use the correct power supply. The power adaptor specifications appear on the serial number label (bottom of the instrument) and in the "Technical specifications" section of this user guide (page 48). Damage due to an incompatible power adaptor is not covered by warranty.



CAUTION! Never disassemble or service the instrument yourself. Do not remove any covers or parts that require the use of a tool to obtain access to moving parts. Operators must be trained before being allowed to perform the hazardous operation. Unauthorized repairs may damage the instrument or alter its functionality, which may void your warranty. Contact your local distributor to arrange for service.



IMPORTANT! If you have any doubt about the compatibility of decontamination or cleaning agents with parts of the equipment or with material contained in it, contact Technical Support (page 63) or your local distributor for information.

Clean the Countess™ II FL Automated Cell Counter

Introduction

We recommend cleaning the Countess[™] II FL Automated Cell Counter periodically to prevent the buildup of dust and dirt that might reduce its performance and cause contamination.



CAUTION! To avoid electrical shock, always turn off the Counters™ II FL Automated Cell Counter and unplug the power cord before cleaning or decontaminating the instrument.



CAUTION! All biological samples and materials that come into contact with them have the potential to transmit infectious diseases and are considered biohazardous. Follow all applicable local, state/provincial, and/or national regulations. Wear appropriate protective eyewear, clothing, and gloves.



IMPORTANT! Using a cleaning or decontaminating method other than that sp manufacturer may result in damage to the instrument.

Clean the touchscreen

- Wipe the touch-screen of the Countess™ II FL Automated Cell Counter using a soft, lint-free cloth moistened with an LCD cleaning solution. Do not apply excessive force during cleaning. Wipe the touch-screen dry immediately after cleaning.
- Ensure that the cleaning solution does not enter the power button, the power inlet, the slide port, or the USB ports.
- Never pour or spray any liquids directly on the instrument to avoid electrical shock when the instrument is plugged in.
- Do not use abrasive cleaning solutions or material to prevent the touch-screen from getting scratched.

Clean the instrument case

- Wipe the instrument case of the Countess™ II FL Automated Cell Counter using a soft, lint-free cloth moistened with distilled water. Wipe the instrument dry immediately after cleaning.
- Ensure that water or other cleaning solutions do not enter the power button, the power inlet, the slide port, or the USB ports.
- Never pour or spray any liquids directly on the instrument to avoid electrical shock when the instrument is plugged in.

Decontaminate the instrument

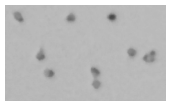
- Wipe the instrument case of the Countess[™] II FL Automated Cell Counter using a soft, lint-free cloth moistened with 70% alcohol. Wipe the instrument dry immediately after cleaning.
- Avoid using a bleach solution, because it may leave a residue of bleach crystals on the instrument.
- Ensure that water or other cleaning solutions do not enter the power button, the power inlet, the slide port, or the USB ports.
- Never pour or spray any liquids directly on the instrument to avoid electrical shock when the instrument is plugged in.

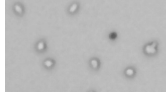
Set nominal focus

Overview

Nominal focus is the Z-point (i.e., depth) around which the auto focus function searches to provide fine focus to the sample.

The auto focus algorithm of the Countess™ II FL Automated Cell Counter is designed to highlight the differences between live and dead cells in the brightfield channel. The optimal focus level is where the "live" cells have a light colored center and the "dead" cells are dark throughout (see examples below).





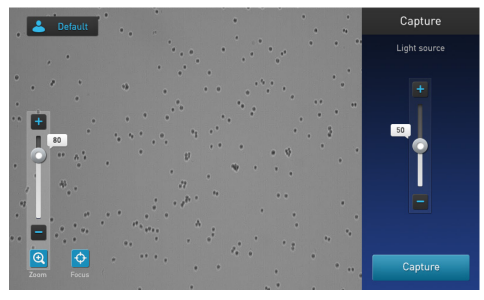
Focus is not optimal

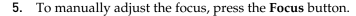
Focus is optimal

To enable optimal auto focus functionality, you may need to initially refine the brightfield focus by adjusting it manually and then setting the nominal focus. This allows the auto focus function to have a set point from which to focus on the cells in subsequent samples.

Set nominal focus

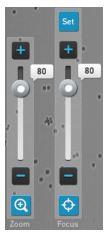
- 1. Prepare the sample by adding $10~\mu L$ of cell suspension to $10~\mu L$ of 0.4% trypan blue stain. Mix the sample mixture well by pipetting up and down a few times.
- 2. Load 10 μ L of the sample mixture into the CountessTM Cell Counting Chamber Slide (page 15) or the CountessTM II FL Reusable Slide (page 18). Let the sample mixture settle for 30 seconds to ensure a uniform focal plane.
- 3. Insert the sample slide into the slide port of the instrument (see page 7), making sure that the side containing the sample is inserted completely.
- 4. When the slide is inserted, the instrument automatically illuminates the sample, sets the intensity of the brightfield light source, and auto focuses on the cells.







6. Use the **Focus slider** or the **plus** and **minus** buttons to refine the brightfield focus.



Note: If needed, Zoom in on the image to adjust focus or lighting.



7. After finding the optimal focus, press **Set** to set the nominal focus. Once the focus has been set, the Set button on the focus slider becomes inactive, confirming that the focus setting has been stored.



Appendix A: Troubleshooting



Note: The software for the CountessTM II FL is updated regularly. If you are having any issues with your experiments, first check the website to see if a new software version is available. You can download the most recent version of the software from **www.thermofisher.com/countessupdate**. You can also register your CountessTM II FL instrument at **www.thermofisher.com/registercountess** to be informed of any future software updates.

Problem	Possible solutions	
Uneven screen illumination (screen is dark on one side, but brighter on the other)	Reset the light cube tray by selecting Change Light Cube on the Instrument Settings screen (page 36).	
Autofocus does not seem to focus on the cells very well	 Make sure there are no bubbles or debris visible on the screen that could interfere with the autofocus and make it more difficult to get the sample in the correct focal plane. Ideally, the live cells should have bright centers compared to the dead cells, which are dark throughout (page 44). 	
the cens very wen	 Setting the nominal focus will improve autofocus consistency with future slides. To set the nominal focus, see page 44. 	
	For cell count and cell viability assays performed in the brightfield, adjust the size, brightness, and circularity gates for both live and dead cells to include all of the cells in the count (page 24).	
Some cells appear on the image but are not included in the count	• For fluorescence assays, adjust the size, brightness, circularity, and fluorescence intensity gates in all available channels to include all of the cells in the count (page 32).	
	• After including all of the cells in the count, you can narrow the count criteria, if you wish to exclude cells of a certain size or certain brightness.	
	• When the gates are fully maximized, the CSV should indicate 0–60 for cell size and 0–255 for brightness.	
Images are very bright and washed out	Enable Auto Lighting from the Profiles menu, or decrease the bright field light intensity before counting the cells.	
Fluorescence is extremely bright and bleeding through into other filters	Decrease the fluorescence light intensity before counting the cells.	
	The beads can settle quickly in solution, which will affect the concentration reading.	
Getting incorrect concentration for the Countess [™] test beads	• Vortex the bead stock on high for a full 30 seconds to resuspend, and add 10 μ L of the bead suspension to 10 μ L of trypan blue without delay.	
	• Pipet the bead and trypan blue mixture up and down several times to make sure it is well mixed, and immediately load $10~\mu L$ into the slide.	

Problem	Possible solutions
	• If you are pipetting different samples from the same cell sample, the variability could be due to pipetting or mixing.
Variable counts for the same sample of cells	 Use recently calibrated pipettors and make sure that the cells are well suspended by pipetting up and down several times before adding trypan blue.
	• Pipet the bead and trypan blue mixture up and down several times to make sure it is well mixed, and load 10 μ L into the slide without delay.
	• If you are counting replicates of the exact same slide, visually inspect that all cells are counted correctly in the image.
Variable counts when performing replicate counts of the same slide	• There may be a slightly different field of view each time a slide is inserted. Depending on the concentration and uniformity of the cells, this will cause some variability when performing replicate counts of the same slide, although it should be less than 10%.
	 When counting fewer cells, a small field of view change for only a small number of cells can have a larger affect. Count cells at a higher concentration to reduce variability.
	• Make sure that you do not shake or agitate the slide between counts.
	 Ensure that the cells are focused correctly so that live cells have bright centers and dead cells are dark throughout (see page 44). If the cells are not well focused and look dark on the screen, the Countess™ II FL will count them as dead cells.
Abnormally high percentage of dead cells or live cells counted as dead	 If cells are well focused, have bright centers, and are being counted as dead, confirm that they are within the appropriate cell size range and try adjusting the settings.
	• If cells are exposed to trypan blue for a long period of time, viability could be affected so slide should be prepared and counted fresh each time for best results.
	• Gate out the debris using the size, brightness, and circularity sliders.
USB drive not recognized by the instrument	• The USB drive must be FAT32 formatted to be recognized by the instrument. If it is not, reformat the USB drive to FAT32 (page 37).
	Try another correctly formatted USB drive.
	 Make sure the USB drive is formatted to FAT32. If it is not, reformat the USB drive to FAT32 before transferring the files onto the USB drive for software update.
Unable to update the Countess™	• The update file must sit on the top level of the USB drive, not within a folder or a subfolder.
software	• File cannot be renamed in any way.
	 File cannot be zipped or compressed during distribution. It must be uncorrupted during transfer and have a .lft suffix.
	If needed, check that the USB port is functional by testing a USB mouse.

Appendix B: Product specifications

Technical specifications

characteristics

Physical Instrument type: Benchtop cell counter and suspension cell-based

assay platform

Instrument dimensions: $9 (W) \times 5\frac{1}{2} (D) \times 9 \text{ inches } (H)$

Weight: 8 lbs

Operating power: 100–240 VAC, 0.58 A MAX

Frequency: 50/60 Hz Electrical input: 12 VDC, 2 A

Installation site: Indoor use only, Class A Environments

(i.e., non-residential or light industrial);

Pollution degree 2.

Operating temperature: 10–40°C

Operating humidity: <80% (non-condensing)

Technical Processing time: ~15 seconds

Sample concentration range: 1×10^4 – 1×10^7 cells/mL

Particle/cell diameter range: 4–60 μm (particles); 7–60 μm (cells)

Required sample volume: 10 µL

Firmware: Countess[™] Automated Cell Counting Platform

Software

USB Drive : 4 Gigabyte

Optics Optics: 3 channels (brightfield and 2 slots for EVOS $^{\text{TM}}$

LED light cubes)

Camera: 5 Mega pixels, 2.5× Optical Magnification

Analysis slide Material: Poly(methyl methacrylate) (PMMA)

Dimensions: $25 \text{ mm (W)} \times 75 \text{ mm (D)} \times 1.7 \text{ mm (H)}$

Chamber volume: 10 μL

EVOS™ light cubes

LED illumination

The Countess™ II FL Automated Cell Counter utilizes an adjustable intensity LED light source provided by the proprietary, user-interchangeable LED light cube (see below). Because the LED light source is as close as possible to the objective, the number of optical elements in the channel is minimized. High-intensity illumination over a short channel increases the efficiency of fluorophore excitation, providing better detection of weak fluorescent signals.

EVOS™ light cubes

Each user-interchangable, auto-configured EVOS™ light cube contains an LED, collimating optics, and filters. In addition to the brightfield channel dedicated to cell count and cell viability assays using Trypan Blue, the Countess™ II FL Automated Cell Counter can accommodate two fluorescent light cubes for multiple-fluorescence research applications.



The table below lists some of the common fluorescent and specialty EVOS™ light cubes available from Thermo Fisher Scientific. For a complete list, go to **www.thermofisher.com/evoslightcubes** or contact Technical Support (see page 63). For instructions on changing the LED light cubes, see page 40.

Light cube	Dye	
DAPI	DAPI, Hoechst™, BFP	
TagBFP	TagBFP	
CFP	ECFP, Lucifer Yellow, Evans Blue	
GFP	GFP, Alexa Fluor™ 488, SYBR™ Green, FITC	
YFP	EYFP, acridine orange + DNA	
RFP	RFP, Alexa Fluor™ 546, Alexa Fluor™ 555, Alexa Fluor™ 568, Cy®3, MitoTracker™ Orange, Rhodamine Red, DsRed	
Texas Red	Texas Red [™] , Alexa Fluor [™] 568, Alexa Fluor [™] 594, MitoTracker [™] Red, mCherry, Cy [™] 3.5	
Cy5	Cy®5, Alexa Fluor™ 647, Alexa Fluor™ 660, DRAQ5™	
Cy5.5	Cy®5.5, Alexa Fluor™ 660, Alexa Fluor™ 680, Alexa Fluor™ 700	
Су7	Cy®7, IRDye 800CW	



Note: The EVOSTM light cubes are available only for the CountessTM II FL Automated Cell Counter. The CountessTM II Automated Cell Counter uses only brightfield illumination and does not support the EVOSTM light cubes.

Appendix C: Ordering information

Countess™ II FL Automated Cell Counter and accessories

The following Countess[™] II FL instruments and instrument accessories are available from Thermo Fisher Scientific. For more information, visit **www.thermofisher.com** or contact Technical Support (page 63).

Product	Quantity	Cat. no.
Countess™ II FL Automated Cell Counter	1 each	AMQAF1000
Countess [™] II power cord with four adapter cords	1 each	AMEP4716
Countess™ II USB drive	1 each	A25751
Countess™ II FL Light Cube Removal Tool	1 each	AMEP4747
Countess™ II FL Disposable Slide Holder	1 each	AMEP4745
Countess™ II FL Reusable Slide Holder	1 each	AMEP4746

Accessory products

The following products can be used with the Countess[™] II FL Automated Cell Counter and are available separately from Thermo Fisher Scientific. For more information, visit **www.thermofisher.com** or contact Technical Support (page 63).

Product	Quantity	Cat. no.
Countess™ Cell Counting Chamber Slides, 50 Slides (100 counts)	1 box*	C10228
Countess™ Cell Counting Chamber Slides, 500 Slides (1000 Counts)	10 boxes*	C10312
Countess™ Cell Counting Chamber Slides, 1250 Slides (2500 Counts)	25 boxes*	C10313
Countess [™] Cell Counting Chamber Slides, 2500 Slides (5000 Counts)	50 boxes*	C10314
Countess™ Cell Counting Chamber Slides, 5000 Slides (10,000 Counts)	100 boxes*	C10315
Countess [™] II FL Reusable Slide	1 each	A25750
Countess [™] Test Beads (1 × 10 ⁶ beads/mL)	1 mL	C10284
Trypan blue stain (0.4 %)	2 × 1 mL	T10282

^{*} Each box of CountessTM Cell Counting Chamber Slides contains 50 slides and 2×1 mL vials of trypan blue (0.4%), sufficient for 100 counts.

Appendix D: CSV file format

CSV file format, explained

Overview

A comma-separated values (CSV) file stores tabular data (numbers and text) in plaintext form. Plain text means that the file is a sequence of characters, with no data that has to be interpreted as binary numbers. A CSV file can be opened with any third party software or spreadsheet program. The table below describes the categories of the Countess™ II data saved as a CSV file and opened with a spreadsheet program.

Category	Column	Name	Description
General	А	Number	Sequential sample run number
	В	File Name	Name of file
	С	Date & Time	Date and time of sample run
	D	Mode	BF-Brightfield or FL-Fluorescence
Trypan	Е	Total Concentration	Concentration of the entire sample
Blue/Brightfield	F	Total cells counted	Total number of cells counted in the sample
	G	Live concentration	Concentration of just the "live" portion of the sample
	Н	Live cells counted	Total number of "live" cells counted
	I	Dead concentration	Concentration of just the "dead" portion of the sample
	J	Dead cells counted	Total number of "dead" cells counted
	K	Viability (%)	Percent viability of the sample based on trypan blue staining
	L	Average size (um)	Average cell size in microns
Fluorescence	М	Cube 1 name	EVOS light cube name in the first (top) position
	N	Cube 1 concentration	Concentration of cells showing fluorescence in the first cube position
	0	Cube 1 (%)	Percentage of the total cells in brightfield that show fluorescence in the first cube position
	Р	Cube 1 cells counted	Total number of cells counted in the first cube position
	Q	Cube 2 name	EVOS light cube name in the second (bottom) position
	R	Cube 2 concentration	Concentration of cells showing fluorescence in the second cube position
	S	Cube 2 (%)	Percentage of the total cells in brightfield that show fluorescence in the second cube position
	Т	Cube 2 cells counted	Total number of cells counted in the second cube position
	U	Cube 1+2 concentration	Concentration of cells showing fluorescence in the first and second cube positions combined
	V	Cube 1+2 (%)	Percentage of the total cells in brightfield that show fluorescence in the first and second cube position combined
	W	Cube 1+2 cells counted	Total number of cells counted in the first and second cube position combined

Category	Column	Name	Description
General Details	Х	Focus value	Focal position number
	Υ	BF Light intensity	Brightfield light intensity value from 0-100%
Trypan	Z	Live Size min	Minimum size of "live" cells in microns
Blue/Brightfield Count Parameters	AA	Live Size max	Maximum size of "live" cells in microns
Count Parameters	AB	Live Brightness min	"Live" adjustment slider value for minimum brightness
	AC	Live Brightness max	"Live" adjustment slider value for maximum brightness
	AD	Live Circularity	"Live" adjustment slider value for circularity
	AE	Dead Size min	Minimum size of "dead" cells in microns
	AF	Dead Size max	Maximum size of "dead" cells in microns
	AG	Dead Bright min	"Dead" adjustment slider value for minimum brightness
	АН	Dead Bright max	"Dead" adjustment slider value for maximum brightness
	Al	Dead Circ	"Dead" adjustment slider value for circularity
Fluorescence	AJ	Cube 1 Light intensity	First (top) light cube light intensity value from 0-100%
Count Parameters	AK	Cube 2 Light intensity	Second (bottom) light cube light intensity value from 0-100%
	AL	BF Size min	Minimum size of "Brightfield" cells in microns
	AM	BF Size max	Maximum size of "Brightfield" cells in microns
	AN	BF Brightness min	"Brightfield" adjustment slider value for minimum brightness
	Α0	BF Brightness max	"Brightfield" adjustment slider value for maximum brightness
	AP	BF Circularity	"Brightfield" adjustment slider value for circularity
	AQ	Cube 1 Brightness min	First (top) light cube adjustment slider value for minimum brightness
	AR	Cube 1 Brightness max	First (top) light cube adjustment slider value for maximum brightness
	AS	Cube 2 Brightness min	Second (bottom) light cube adjustment slider value for minimum brightness
	АТ	Cube 2 Brightness max	Second (bottom) light cube adjustment slider value for maximum brightness

Appendix E: Safety

Safety conventions used in this document

Safety alert words

Four safety alert words appear in this document at points where you need to be aware of relevant hazards. Each alert word–**IMPORTANT**, **CAUTION**, **WARNING**, **DANGER**–implies a particular level of observation or action:



IMPORTANT! Provides information that is necessary for proper instrument operation, accurate installation, or safe use of a chemical.



CAUTION! – Indicates a potentially hazardous situation that, if not avoided, may result in minor or moderate injury. It may also be used to alert against unsafe practices.



WARNING! – Indicates a potentially hazardous situation that, if not avoided, could result in death or serious injury.



DANGER! – Indicates an imminently hazardous situation that, if not avoided, will result in death or serious injury. This signal word is to be limited to the most extreme situations.

Except for **IMPORTANT**! safety alerts, each safety alert word in the document appears with an open triangle figure that contains a hazard symbol. These hazard symbols are identical to the hazard icons that are affixed to the instrument (see "Safety symbols").

Symbols on instruments

Electrical symbols

The following table describes the electrical symbols that may be displayed.

Symbol	Description
	Indicates the On position of the main power switch.
0	Indicates the Off position of the main power switch.
ψ	Indicates a standby switch by which the instrument is switched on to the Standby condition. Hazardous voltage may be present if this switch is on standby.
Φ	Indicates the On/Off position of a push-push main power switch.
÷	Indicates a terminal that may be connected to the signal ground reference of another instrument. This is not a protected ground terminal.
	Indicates a protective grounding terminal that must be connected to earth ground before any other electrical connections are made to the instrument.
~	Indicates a terminal that can receive or supply alternating current or voltage.
=	Indicates a terminal that can receive or supply alternating or direct current or voltage.

Safety symbols

The following table describes the safety symbols that may be displayed. Each symbol may appear by itself or in combination with text that explains the relevant hazard (see "Safety labels on instruments"). These safety symbols may also appear next to DANGERS, WARNINGS, and CAUTIONS that occur in the text of this and other product-support documents.

Symbol	Description		
<u>^</u>	Indicates that you should consult the manual for further information and to proceed with appropriate caution.		
4	Indicates the presence of an electrical shock hazard and to proceed with appropriate caution.		
<u>M</u>	Indicates the presence of a hot surface or other high-temperature hazard and to proceed with appropriate caution.		
*	Indicates the presence of a laser inside the instrument and to proceed with appropriate caution.		
	Indicates the presence of moving parts and to proceed with appropriate caution.		
	Indicates the presence of a biological hazard and to proceed with appropriate caution.		
	Indicates the presence of an ultraviolet light and to proceed with appropriate caution.		

Environmental symbols

The following symbol applies to all Thermo Fisher Scientific electrical and electronic products placed on the European market after August 13, 2005.

Symbol	Description		
	Do not dispose of this product as unsorted municipal waste. Follow local municipal waste ordinances for proper disposal provisions to reduce the environmental impact of waste electrical and electronic equipment (WEEE).		
∕ • \	European Union customers:		
	Call your Customer Service representative for equipment pick-up and recycling. See www.thermofisher.com for a list of customer service offices in the European Union.		

Safety labels on instruments

The following CAUTION, WARNING, and DANGER statements may be displayed Thermo Fisher Scientific instruments in combination with the safety symbols described in the preceding section.

Hazard symbol	English	Français
<u></u>	CAUTION! Hazardous chemicals. Read the Safety Data Sheets (SDSs) before handling.	ATTENTION! Produits chimiques dangereux. Lire les fiches techniques de sûreté de matériels avant toute manipulation de produits.
	CAUTION! Hazardous waste. Refer to SDS(s) and local regulations for handling and disposal.	ATTENTION! Déchets dangereux. Lire les fiches techniques de sûreté de matériels et la régulation locale associées à la manipulation et l'élimination des déchets.
<u></u>	DANGER! High voltage.	DANGER! Haute tension.
<u> </u>	WARNING! To reduce the chance of electrical shock, do not remove covers that require tool access. No user-serviceable parts are inside. Refer servicing to Thermo Fisher Scientific qualified service personnel.	AVERTISSEMENT! Pour éviter les risques d'électrocution, ne pas retirer les capots dont l'ouverture nécessite l'utilisation d'outils. L'instrument ne contient aucune pièce réparable par l'utilisateur. Toute intervention doit être effectuée par le personnel de service qualifié venant de Thermo Fisher Scientific.
*	DANGER! Class 3B visible and/or invisible laser radiation present when open. Avoid exposure to beam.	DANGER! Rayonnement visible ou invisible d'un faisceau laser de Classe 3B en cas d'ouverture. Evitez toute exposition au faisceau.
	CAUTION! Moving parts. Crush/pinch hazard.	ATTENTION! Pièces en mouvement, risque de pincement et/ou d'écrasement.

General instrument safety



WARNING! PHYSICAL INJURY HAZARD. Use this product only as specified in this document. Using this instrument in a manner not specified by Thermo Fisher Scientific may result in personal injury or damage to the instrument.

Operating the instrument

Ensure that anyone who operates the instrument has:

- Received instructions in both general safety practices for laboratories and specific safety practices for the instrument.
- Read and understood all applicable Safety Data Sheets (SDSs). See "Documentation and support" (page 63).

Safety precautions

- Do not install the instrument in heavy humidity such as a greenhouse or an
 incubator to avoid a danger of electric shock. If water or other material enters
 the instrument, the adaptor, or power inlet, disconnect the power cord and
 contact a service person. For operating environment, refer to "Operating
 environment" (page 8).
- Do not press the main plug or power cord with wet hands.
- Always ensure that the power supply input voltage matches the voltage available in your location.
- Do not install the instrument on a slant or a place prone to vibrations, which induces the risk of instrument malfunction or damage of the instrument.
- Never insert any objects into the air vents of the instrument as this could result in electrical shock, personal injury, and equipment damage.
- Plug the power cord firmly into the wall outlet and the instrument.
- To avoid potential shock hazard, make sure that the power cord is properly grounded.
- Be sure to position the equipment such that it is easy to disconnect the instrument.
- Turn off the instrument before unplugging the power cord and/or moving the instrument.
- If the instrument is broken or dropped, disconnect the power cord and contact a service person. Do not disassemble the instrument.
- Use only authorized accessories (adaptor, power cord, and USB drive).
- If the instrument emits smoke, disconnect the power cord from the wall outlet and contact a service person.

Cleaning or decontaminating the instrument



CAUTION! Using cleaning or decontamination methods other than those recommended by the manufacturer may compromise the safety or quality of the instrument.

Removing covers or parts of the instrument



CAUTION! PHYSICAL INJURY HAZARD. The instrument is to be serviced only by trained personnel or vendor specified in the user guide.

Chemical safety

Chemical hazard warning



WARNING! CHEMICAL HAZARD. Before handling any chemicals, refer to the Safety Data Sheet (SDS) provided by the manufacturer, and observe all relevant precautions.



WARNING! CHEMICAL STORAGE HAZARD. Never collect or store waste in a glass container because of the risk of breaking or shattering. Reagent and waste bottles can crack and leak. Each waste bottle should be secured in a low-density polyethylene safety container with the cover fastened and the handles locked in the upright position. Wear appropriate eyewear, clothing, and gloves when handling reagent and waste bottles.

General safety guidelines

To minimize the hazards of chemicals:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the SDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open.
 Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the SDS.
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the SDS.
- Comply with all local, state/provincial, or national laws and regulations related to chemical storage, handling, and disposal.

Chemical waste safety

Chemical waste hazard



CAUTION! HAZARDOUS WASTE. Refer to Safety Data Sheets (SDSs) and local regulations for handling and disposal.

Chemical waste safety guidelines

To minimize the hazards of chemical waste:

- Read and understand the Safety Data Sheets (SDSs) provided by the manufacturers of the chemicals in the waste container before you store, handle, or dispose of chemical waste.
- Provide primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the SDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the SDS.
- Handle chemical wastes in a fume hood.
- After emptying the waste container, seal it with the cap provided.
- Dispose of the contents of the waste tray and waste bottle in accordance with good laboratory practices and local, state/provincial, or national environmental and health regulations.

Waste disposal

If potentially hazardous waste is generated when you operate the instrument, you must:

- Characterize (by analysis, if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure the health and safety of all personnel in your laboratory.
- Ensure that the instrument waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.



IMPORTANT! Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

Electrical safety



DANGER! ELECTRICAL SHOCK HAZARD. Severe electrical shock can result from operating the Countess™ II FL Automated Cell Counter without its instrument panels in place. Do not remove instrument panels. High-voltage contacts are exposed when instrument panels are removed from the instrument.

Fuses



WARNING! FIRE HAZARD. For continued protection against the risk of fire, replace fuses only with fuses of the type and rating specified for the instrument.

Power



DANGER! ELECTRICAL HAZARD. Grounding circuit continuity is vital for the safe operation of equipment. Never operate equipment with the grounding conductor disconnected.



DANGER! ELECTRICAL HAZARD. Use properly configured and approved line cords for the voltage supply in your facility.



DANGER! ELECTRICAL HAZARD. Plug the system into a properly grounded receptacle with adequate current capacity.

Overvoltage rating

The Countess™ II FL Automated Cell Counter has an installation (overvoltage) category of II, and is classified as portable equipment.

Biological hazard safety



WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, and blood of humans and other animals have the potential to transmit infectious diseases. Follow all applicable local, state/provincial, and/or national regulations. Wear appropriate protective eyewear, clothing, and gloves. Read and follow the guidelines in these publications.

ATTENTION! BIOHAZARD. Les échantillons biologiques tels que les tissus, les fluides corporels et le sang des humains et d'autres animaux ont la possibilité de transmettre des maladies infectieuses. Suivre tous les règlements municipaux, provinciaux/provincial et / ou nationales en vigueur. Porter des lunettes de protection approprié, des vêtements et des gants.

In the U.S.:

- U.S. Department of Health and Human Services guidelines published in Biosafety in Microbiological and Biomedical Laboratories (stock no. 017-040-00547-4;
 - www.cdc.gov/OD/ohs/biosfty/bmbl4/bmbl4toc.htm)
- Occupational Safety and Health Standards, Bloodborne Pathogens (29 CFR§1910.1030;
 - www.access.gpo.gov/nara/cfr/waisidx_01/29cfr1910a_01.html)
- Your company's/institution's Biosafety Program protocols for working with/handling potentially infectious materials.
- Additional information about biohazard guidelines is available at: www.cdc.gov

In the EU:

- Check your local guidelines and legislation on biohazard and biosafety precaution, and the best practices published in the World Health Organisation (WHO) Laboratory Biosafety Manual, third edition
 - www.who.int/csr/resources/publications/biosafety/WHO_CDS_CSR_LYO_2004_11/en/

Safety and electromagnetic compatibility (EMC) standards

This section provides information on:

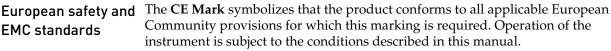
- U.S. and Canadian safety standards
- European safety and EMC standards
- Australian EMC standards

U.S. and Canadian safety standards



The CSA C/US Mark signifies that the product meets applicable U.S. and Canadian standards, including those from CSA, CSA America, ANSI, ASME, ASSE, ASTM, NSF and UL.

EMC standards





The protection provided by the instrument may be impaired if the instrument is used in a manner not specified by Thermo Fisher Scientific.

Australian EMC standards



The C-Tick Mark indicates conformity with Australian and New Zealand standards for electromagnetic compatibility.

Documentation and support

Obtaining support

Technical support

Visit **www.thermofisher.com/support** for the latest in services and support, including:

- Worldwide contact telephone numbers
- Product support, including:
 - Product FAQs
 - Software, patches, and updates
- Order and web support
- Product documentation, including:
 - User guides, manuals, and protocols
 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

Life Technologies Corporation and/or its affiliate(s) warrant their products as set forth in the Life Technologies' General Terms and Conditions of Sale found on Life Technologies' website at www.lifetechnologies.com/termsandconditions.

If you have any questions, please contact Life Technologies at www.lifetechnologies.com/support.

