Fluorescence-based fixative and vital staining of lipid droplets in Caenorhabditis elegans reveal fat stores using microscopy and flow cytometry approaches

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Abstract The proportions of body fat and fat-free mass are determining factors of adiposity-associated diseases. Work in Caenorhabditis elegans has revealed evolutionarily conserved pathways of fat metabolism. Nevertheless, analysis of body composition and fat distribution in the nematodes has only been partially unraveled because of methodological difficulties. We characterized metabolic C. elegans mutants by using novel and feasible BODIPY 493/503-based fat staining and flow cytometry approaches. Fixative as well as vital BODIPY staining procedures visualize major fat stores, preserve native lipid droplet morphology, and allow quantification of fat content per body volume of individual worms. Colocalization studies using coherent anti-Stokes Raman scattering microscopy, Raman microspectroscopy, and imaging of lysosome-related organelles as well as biochemical measurement confirm our approaches. We found that the fat-to-volume ratio of dietary restriction, TGF-β, and germline mutants are specific for each strain. In contrast, the proportion of fat-free mass is constant between the mutants, although their volumes differ by a factor of 3.4. Our approaches enable sensitive, accurate, and high-throughput assessment of adiposity in large C. elegans populations at a single-worm level.—Klapper, M., M. Ehmke, D. Palgunow, M. Böhme, C. Matthäus, G. Bergner, B. Dietzkek, J. Popp, and F. Döring. Fluorescence-based fixative and vital staining of lipid droplets in Caenorhabditis elegans reveal fat stores using microscopy and flow cytometry approaches. J. Lipid Res. 2011. 52: 1281–1293.

Supplementary key words C. elegans • CARS microscopy • fatty acid • fatty acid metabolism • fluorescence microscopy • obesity • triglycerides

Body composition with respect to white adipose tissue and lean body mass is of central importance and a determining factor of adiposity-associated perturbations, such as diabetes type 2, cardiovascular disease, and cancer. Therefore, modulation of body composition and especially the reduction of body fat are the main foci of dietary and lifestyle interventions. In the last few years, Caenorhabditis elegans became a model organism for investigating fat storage and metabolism, as reviewed by different authors (1–3). Based on forward genetics (4–6), functional genomics, and candidate gene approaches (7, 8), more than 400 genes possibly involved in fat storage were described in C. elegans. Most of these genes are evolutionarily conserved and allocated in common pathways and are of relevance for the study of human adiposity.

Although C. elegans was extensively used as a model organism to study the genetic and functional basis of fat storage, robust lipid droplet markers such as perilipin are not available at the moment (2). Appropriate methods concerning the distribution behavior of dyes in the worm or the reproducibility of staining procedures are still under discussion (3). The classical fixative Sudan black dye (9) is shown to be very error prone in C. elegans because of the last ethanol-based washing steps. Recently O’Rourke et al. (10) demonstrated that the vital dyes Nile red and C1-C12-BODIPY-labeled fatty acid, which were used in more than 75 reports, stain lysosome-like granules rather than lipid droplets within the C. elegans intestine. These dyes were shown not to localize with the fixative neutral fat fluorophore LipidTox (10) and with the label-free lipid droplet visualizing coherent anti-Stokes Raman scattering (CARS).

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Abbreviations: BP, bandpass; CARS, coherent anti-Stokes Raman scattering; COPAS, cytometry-based object parametric analysis and sorting system; LRO, lysosome-related organelle; OPO, optical parametric oscillator; TOF, time of flight; TAG, triacylglycerol.

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of dietary restriction, TGF-β content and body volume revealed specific fat proportions and a novel quantitative BODIPY 493/503-based flow cytometry method allowing parallel measurement of fat and LRO markers in C. elegans (13). Here, we have developed a protocol for staining and quantification of fixed worms, using BODIPY 493/503. Furthermore, body composition analysis and a novel quantitative BODIPY 493/503-based flow cytometry method allowing parallel measurement of fat content and body volume revealed specific fat proportions of dietary restriction, TGF-β, and germline mutants at single-worm level. Additionally, we show that vital staining of worms with BODIPY 493/503 using a short incubation time leads to a very efficient staining of lipid droplets. This method may be suitable for screening assays.

**MATERIALS AND METHODS**

**Strains and C. elegans maintenance**

Bristol N2 was used as the wild-type strain. The glp-1 (e2141), eat-2 (ad465), anddaf-7 (e1372) mutant strains were used. The Glp-1 mutant was a gift from Ralf Schnabel (Technical University of Braunschweig, Germany). Other strains were obtained from the Caenorhabditis Genetic Center. Nematodes were cultured on Escherichia coli OP50 lawns on NGM agar plates, as described previously (22), at 15°C, 20°C, or 25°C. For all experiments, strains were synchronized by hypochlorite treatment of gravid adults.

**BODIPY 493/503 staining and quantification**

The BODIPY 493/503 stock solution (1 mg/ml) was prepared in DMSO. A fresh solution of BODIPY 493/503 diluted in M9 buffer at the indicated concentrations was used for staining. For fixative staining, freshly harvested nematodes were washed in M9 buffer, incubated in 4% paraformaldehyde solution for 15 min, which was followed by three freeze/thaw cycles in liquid nitrogen. After paraformaldehyde solution was removed by three washing steps with M9, worms were incubated in a volume of 500 µl of 1 µg/ml BODIPY 493/503 (Invitrogen) in M9 for 1 h at room temperature. Nematodes were washed three times with M9 buffer and were used for microscopy or flow cytometry analysis. For vital staining, freshly harvested nematodes were incubated in M9/BODIPY 493/503 (6.7 µg/ml) solution for 20 min. An incubation of 20 min generated a sufficient BODIPY 493/503 signal and prevented starvation of the worms and staining of LROs. Afterward, worms were washed three times with M9 and used immediately for further experiments. BODIPY 493/503 fluorescence was visualized using an Axio Observer D1 inverted microscope (Zeiss) using filter 38HE (excitation, bandpass [BP] 470/40; beam splitter FT 495; emission BP 525/50). Images were captured with an AxioCam MRm camera. Quantification of fixative BODIPY 493/503 staining of 20–30 whole worms/experiment was performed using Axio-View software version 4.8 (Zeiss).

Additionally, fluorescence was quantified using the cytometry-based object parametric analysis and sorting system (COPAS), Biosort (Union Biotometrica), which measures time of flight (TOF; nematode length), decrease in laser light when a worm passes through the laser beam (extinction), and green fluorescence (BODIPY fluorescence) for each animal. The setup was equipped with a 488 nm multilane argon laser. Green and red fluorescence emissions were measured using a filter of 510 nm, with a bandwidth of 23 nm. Worms were washed from plates, placed in the COPAS sample cup, and analyzed. COPAS settings were as follows: gains: extinction (EXT), 1; green, 1; yellow, 1 and red, 1; thresholds: signal, 50 and TOF minimum, 100. Voltages of photomultiplier tubes for the yellow, green, and red channels were set to 600 V. Only objects with TOF values of adult nematodes and signals detected in the green channel were included in quantification data. The background fluorescence of unstained worms was subtracted in microscopy and COPAS analyses. Nematodes (250 to 2,000) in each of two to three independent experiments were analyzed using COPAS. Microscopy and COPAS settings were kept identical to allow a direct comparison between groups.

For colocalization studies, nematodes were stained with 1 µg/ml BODIPY 493/503 and LipidTOX deep red (1:200 dilution; Invitrogen) or Nile red (1 µg/ml; Sigma), for 1 h. Confocal images were taken using a Leica TCS SP1 confocal system (Leica) equipped with an HCX PL APO CS 63.0 × 1.32 oil-immersion objective. BODIPY 493/503 and LipidTOX deep red were excited at 488 and 633 nm, respectively. Signals were collected with a TD488/543/633 beam splitter at 500–565 nm and 640–720 nm, respectively. BODIPY 493/503 and Nile red were excited at 488 nm. Signals were collected with a TD488/543/633 beam splitter at 500–520 nm and 600–650 nm, respectively. Autofluorescence and LysoTracker red in vital worms were detected using filter sets 49 (excitation, G 665; beam splitter, G 395; emission, BP 445/50) (23) and 43 HE (excitation BP 550/25; beam splitter, FT 570; emission, BP 605/70), respectively. At all fluorophore combinations, an optimal separation of channels was ensured. The quantitative colocalization analysis was performed using ImageJ plugin JACoP (Just Another Colocalization Plugin) (24).

**Oil Red O staining**

After 15 min of 4% paraformaldehyde fixation followed by three freeze/thaw cycles in liquid nitrogen, worms were washed with PBS and dehydrated in 60% isopropanol. Afterward, worms were stained in filtered Oil Red O staining solution (60% Oil Red O stock solution [5 mg/ml isopropanol]/40% water). Worms were washed with PBS and observed with an Axio Observer D1 inverted microscope (Zeiss). Images were captured with an AxioCamICc1 camera.

**Flow cytometry and biochemical assays**

Synchronized worms were cultivated until they reached adulthood. Synchronized adult nematodes (300–600) were counted for each biological replicate by using the COPAS Biosort system. The worms were resuspended in 70 µl of lysis buffer (50 mM Tris, pH 7.5; 150 mM NaCl; 1 mM EDTA, pH 8.0; 0.5% NP-40; protease inhibitors) and sonicated using a Sonopuls HD 2070 (Bandelin) with an MS72 sonotrode at 30% power for 2 × 8 s. Insoluble material was removed by centrifugation at 21,000 g for 20 min. We assayed triacylglycerol (TAG) content spectrophotometrically, using an enzymatic assay kit (Analyticon) with a TAG standard (Biovision) according to the manufacturer’s instructions. As shown in supplementary Fig. III, the TAG assay kit is suitable for determination of TAG in C. elegans. The protein content was determined using a bicinchoninic acid reagent kit (Pierce) with threshold: signal, 50 and TOF minimum, 100. Voltages of photomultiplier tubes for the yellow, green, and red channels were set to 600 V. Only objects with TOF values of adult nematodes and signals detected in the green channel were included in quantification data. The background fluorescence of unstained worms was subtracted in microscopy and COPAS analyses. Nematodes (250 to 2,000) in each of two to three independent experiments were analyzed using COPAS. Microscopy and COPAS settings were kept identical to allow a direct comparison between groups.

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BSA as standard. We performed two to three independent experiments for each genotype and condition with two replicates, respectively. We measured total TAG and protein contents for each homogenate in duplicate in two independent measurements, respectively. To account for differences in total TAG content per worm, we used protein and body volumes as covariates in the analysis of the data.

CARS microscopy

Worms were washed in M9 buffer, resuspended in 2% NaN₃ and placed on glass slide. A cover glass was then gently applied to the sample. CARS images shown in Fig. 3 were recorded using a CARS microscope that was custom built by A. Zumbusch and M. Winterhalder at the University of Konstanz, based on a Leica TCS SP5 multiphoton microscope. As excitation source for the pump beam, the output of an optical parametric amplifier (APE, Berlin, Germany) was used, whereas the output of a Ti:sapphire laser (Coherent Mira, Santa Clara, CA) served as the Stokes beam. Both sources were electronically synchronized (coherent SynchroLock, Santa Clara, CA). Images were recorded at a laser frequency difference of 2.845 cm⁻¹, corresponding to an aliphatic CH vibrational resonance.

The setup used for CARS microscopy shown in Fig. 4, 5, and 10 was described previously (25). For the experiments, narrowband (bandwidth, 0.5–1 nm) ps pulses were used to match the C-H stretching region at about 2,900 cm⁻¹. The Stokes pulses were provided by a Ti:sapphire laser centered at 831.7 nm. In addition, this laser served as a pump source for an optical parametric oscillator (OPO), which was tuned to 670 nm for resonant and 678 nm for vibrationally nonresonant excitation. These last settings were used as controls to ensure that image contrast stemmed from a CARS process. After the pump and Stokes pulses were overlapped both temporally and spatially by a delay stage and a longpass filter, the two laser beams were directed into a commercial laser scanning microscope (510 Meta; Zeiss). The two beams were focused onto the object plane by a 63×/NA 1.40 oil immersion objective, which was used for CARS and fluorescence measurements as well. The power of the Stokes pulse was about 12 mW, and the power of the pump pulse was about 16 mW at the sample. The anti-Stokes signal at 561 nm was detected in forward direction and spectrally filtered from the incoming beams by the combination of shortpass and bandpass filters and detected by a photomultiplier. For every sample position, a depth profile (z-stack) was measured, recording 30 focal plane images separated by 1 µm each. The image size was 512 × 512 pixels, and the image acquisition time per image plane was about 6.3 s.

Two-photon fluorescence

The two-photon fluorescent signal was recorded simultaneously with the CARS images. The signal, which was generated by an air-cooled argon-ion laser (series 77; Lasos, Jena, Germany). The exciting laser radiation is coupled to a Zeiss microscope through a wavelength-specific single-mode optical fiber. The incident laser beam is collimated via an achromatic lens and passes through a holographic BP filter before it is focused onto the sample through the microscope objective. A Nikon Fluor (60×/1.00 NA, WD = 2.0 mm) water immersion objective was used in the studies reported here. The sample is located on a piezo-electrically driven microscope scanning stage with an x-y resolution of approximately 3 nm and a repeatability of ±1 nm, and a z resolution of approximately 0.5 nm and ±2 nm repeatability. The sample is scanned through the laser focus in a raster pattern at a constant stage speed of fractions of a micrometer per second. The continuous motion prevents sample degradation at the focal point of the laser beam. Spectra are collected at a 0.6 µm grid, with a dwell time of 0.25 s. Preceding fluorescent images were recorded at approximately 0.1 mW and an illumination time of 0.1 s. Raman images were reconstructed using a spectral decomposition algorithm based on vertex component analysis (26). Fluorescence images are based on fluorescence intensities.

RESULTS

A BODIPY 493/503-based protocol stains major fat stores in C. elegans and preserves native lipid droplet morphology

In order to characterize fat stores in C. elegans, we developed a gentle staining procedure with short fixation times and without dehydration steps, using the lipid droplet-specific fluorophore BODIPY 493/503. As shown in Fig. 1A and B, the BODIPY 493/503-based protocol preserves worm anatomy and stains spherical structures in the intestine and hypodermis. These compartments are described as the major fat stores of C. elegans wild-type N2 (2). Droplets in oocytes and embryos are also stained by BODIPY 493/503. The BODIPY 493/503-labeled intracellular vesicles are fine and distinctly distributed (Fig. 1C, D). Most of the droplets are <3 µm in diameter. To provide evidence that the BODIPY 493/503-labeled vesicles are lipid droplets, we tested our protocol with LipidTox deep red and Nile red and performed colocalization studies. As shown in Fig. 2, BODIPY 493/503 stains the same compartments and droplets that the two other neutral lipid dyes do. To measure the extent of overlap between the BODIPY 493/503 stain and that of the two other dyes, we calculated the Pearson correlation coefficient (r). The value of this coefficient ranges from −1 to 1; the higher the value of r, the stronger the correlation between the red and green images. The mean Pearson correlation coefficients ± standard deviations (SD) for BODIPY 493/503 with LipidTox, r = 0.927 ± 0.0056, and Nile red, r = 0.934 ± 0.027 (3–5 stacks), reported a high degree of colocalization of BODIPY 493/503 with LipidTox deep red and Nile red. We further proved our BODIPY 493/503 staining procedure with CARS microscopy, a nonlabeling method for visualizing lipid droplets in C. elegans in vivo without fixation steps (11, 27) or after fixation with paraformaldehyde (28). As shown in Fig. 3, CARS microscopy of vital worms shows intestinal and hypodermal lipid droplets similar in size, morphology, and distribution to droplets in images obtained by fixative
Quantification of BODIPY 493/503-based microscopy
data agreed with biochemical analysis and revealed low
fat content of eat-2 mutants and high fat levels of daf-7
and glp-1 mutants

The fat content of many C. elegans mutants is still under
discussion because the well-established vital dyes Nile red
and C1-C12-BODIPY-labeled fatty acids are not accurate
proxies of fat stores in the worm (10). Here we selected
the eat-2(ad465), daf-7(e1372), and glp-1(e2141) strains as
principal fat storage mutants in order to quantify their fat
content with BODIPY 493/503. The eat-2(ad465) mutant,
a genetic model of dietary restriction, lacks a functional

Fig. 1. BODIPY 493/503 staining protocol preserves worm anatomy and stains neutral lipids in the major storage compartments, intestine and hypodermis, and in oocytes and embryos. Brightfield (A) and BODIPY 493/503 confocal images (B) of 1-day-old adult wild-type worms are shown in 3D projection of ~55 images from a z-stack at 1 µm intervals. Five segments from a wild-type worm were imaged with a 63× oil immersion objective. The anterior part is on the left. Bar, 50 µm. (C) Magnified view of hypodermal lipid droplets in head. (D) Magnified view of lipid droplets in proximal intestine. Bar, 10 µm.

BODIPY 493/503 staining. Furthermore, colocalization
experiments revealed that BODIPY 493/503 stains the
same vesicles that are detected with CARS microscopy
(Fig. 4). As a second method that allows a label-free visu-
alization of lipids, proteins, and other metabolites, we
applied Raman microspectroscopy of fixed BODIPY
493/503-stained animals. As shown in Fig. 5, the BODIPY
493/503-labeled structures overlap with lipid signals of
Raman spectra, and similar signal intensities are observed.
Thus, our data demonstrate that the BODIPY 493/503-
stained organelles in fixed animals are lipid droplets with
a well-preserved morphology.

Fig. 2. BODIPY 493/503 signals colocalize with the
dyes Nile red and LipidTox deep red. Fixed worms were stained with BODIPY 493/503 and LipidTox deep red (A–C) or with BODIPY 493/503 and Nile red (D–F). Images were taken of proximal intestine (A, D), middle intestine with uterus (B, E), and distal intestine with tail hypodermis (C, F). A, anterior; P, posterior; I, intestine; G, gonad; E, embryo, H, hypodermis. Bar, 50 µm; 10 µm in magni-

ified views.
rather than the absolute fat content of an organism, the fat-to-fat-free ratio is one of the most relevant physiological parameters used to evaluate body composition and adiposity (35). To calculate this ratio in the eat-2, daf-7, and glp-1 mutants, we also determined the protein fraction. A part of the protein fraction contributes to the metabolic active body compartment of the nematodes. As shown in Fig. 7B, the protein content per worm of the eat-2 (96.2 ± 7.4 ng/worm) mutant is 2.4-fold lower and that of the daf-7 (292.7 ± 6.6 ng/worm) mutant is 30% higher than in the wild type (227 ± 7.1 ng/worm). The glp-1 and N2 strains differ slightly in protein content at permissive as well as at restrictive temperatures. The TAG and protein values were used in order to calculate fat-to-fat-free ratios. Based on these values, the eat-2 mutant could be described as a lean mutant, whereas the daf-7 and glp-1 mutants are obese (Fig. 7C). Another central parameter for evaluating lean-ness or adiposity is the proportion of protein or fat in relationship to the body volume. Volumes (V) of the eat-2, daf-7, and glp-1 mutants were determined using a worm-adapted cylinder volume formula that includes worm area (A) and perimeter (P) (36, 37).

\[ V = \pi \left( P + \sqrt{P^2 - 16A} \right) \left( P - \sqrt{P^2 - 16A} \right)^3 / 256 \]
are the most reliable parameters for comparing the levels of adiposity in different C. elegans strains.

To measure the fat content per volume at the single-worm level in large C. elegans populations, we developed a high-throughput method based on BODIPY 493/503 and the flow cytometry sorting system, COPAS. In principal, the method simultaneously quantifies the integrated fluorescence intensity of BODIPY 493/503 (green fluorescence), the length of the worms (TOF), and the body volume-dependent decrease in laser light (extinction, 488 nm) of single worms. Because fixative BODIPY 493/503 protocol may influence the volume of the worms, we checked this putative confounder. There was no significant change in worm volume when vital and fixed animals were compared (Fig. 8A). The extinction signals of worms obtained by

The body volumes of the different strains vary by a factor of 3 (Fig. 7D). The eat-2 mutant shows a markedly lower body volume (1.08 ± 0.14 nl/worm) than the N2 (2.7 ± 0.08 nl/worm). The volume of the daf-7 mutant is 1.3-fold higher than the wild type. The body volume of the glp-1 and N2 strains are almost identical at permissive temperature (15°C). At restrictive temperature, the glp-1 mutant shows a 20% reduction in body volume in comparison to that of N2. Interestingly, the TAG content per volume (Fig. 7F) is specific for each strain and represents leanness of the eat-2 mutant (64% of N2) and fatness of the daf-7 (150%) and glp-1 (163%) mutants. In contrast, the protein-to-volume ratios representing the proportions of fat-free mass in the worm are constant among strains (Fig. 7E). Taken together, the fat-to-fat-free mass and the fat-to-volume ratio are the most reliable parameters for comparing the levels of adiposity in different C. elegans strains.

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The body volumes of the different strains vary by a factor of 3 (Fig. 7D). The eat-2 mutant shows a markedly lower body volume (1.08 ± 0.14 nl/worm) than the N2 (2.7 ± 0.08 nl/worm). The volume of the daf-7 mutant is 1.3-fold higher than the wild type. The body volume of the glp-1 and N2 strains are almost identical at permissive temperature (15°C). At restrictive temperature, the glp-1 mutant shows a 20% reduction in body volume in comparison to that of N2. Interestingly, the TAG content per volume (Fig. 7F) is specific for each strain and represents leanness of the eat-2 mutant (64% of N2) and fatness of the daf-7 (150%) and glp-1 (163%) mutants. In contrast, the protein-to-volume ratios representing the proportions of fat-free mass in the worm are constant among strains (Fig. 7E). Taken together, the fat-to-fat-free mass and the fat-to-volume ratio are the most reliable parameters for comparing the levels of adiposity in different C. elegans strains.

To measure the fat content per volume at the single-worm level in large C. elegans populations, we developed a high-throughput method based on BODIPY 493/503 and the flow cytometry sorting system, COPAS. In principal, the method simultaneously quantifies the integrated fluorescence intensity of BODIPY 493/503 (green fluorescence), the length of the worms (TOF), and the body volume-dependent decrease in laser light (extinction, 488 nm) of single worms. Because fixative BODIPY 493/503 protocol may influence the volume of the worms, we checked this putative confounder. There was no significant change in worm volume when vital and fixed animals were compared (Fig. 8A). The extinction signals of worms obtained by
COPAS correlated well ($r^2 = 0.81$) with their volumes (Fig. 8C) and were not influenced by fixation (Fig. 8B). Thus, the COPAS-derivative fluorescence signals of BODIPY 493/503-fixed worms can be used as an accurate proxy for the volume of the worms. Next, we measured BODIPY 493/503 fluorescence emission of fat storage mutants by using COPAS. The resulting averaged BODIPY 493/503 signals per worm of the eat-2, daf-7, and glp-2 mutants are in agreement with fat contents obtained by BODIPY 493/503 microscopy and biochemical measurement (Table 1). Finally, BODIPY 493/503 fluorescence and extinction values of single worms were plotted. As shown in Fig. 8D, E, populations of the eat-2, daf-7, and N2 strains show distinct distributions and differences in fat contents and body volumes. To illustrate this for the glp-1 mutant in more detail, N2 worms and glp-1 mutants showed almost identical BODIPY 493/503 signals, body lengths (Fig. 8F), and body vaxedependent extinction signals (Fig. 8H), indicating similar body compositions for both strains at permissive temperature. At restrictive temperature (25°C), the glp-1 mutants showed higher fat content but similar body lengths compared to those of N2 (Fig. 8G). The body volume-dependent extinction signals of the glp-1 strain were slightly lower than those of N2 (Fig. 8I) and are in agreement with direct body volume measurements using image analysis (Fig. 7D). As a final result, the glp-1 mutant showed a 1.6-fold higher fat-to-volume ratio than the N2 strain at restrictive temperature, indicating fattiness of the glp-1 mutant. Thus, data for BODIPY 493/503-based flow cytometry represents the proportion of fat at the single-worm level in large C. elegans populations.

Furthermore, we stained the eat-2, daf-7, and glp-2 mutants with Nile Red after fixation and measured the red fluorescence and extinction by flow cytometry. As shown in supplementary Fig. 1, where red fluorescence signal and extinction are plotted, the relative fluorescence data provide quantitative information about fat content of the mutants in relation to that of the wild type N2. For example, the glp-1 mutant shows a 1.8-fold higher fat-to-volume ratio than the N2. Therefore, fixative Nile red is, as also reported by Brooks et al. (15), a useful quantitative marker for worm TAG content, demonstrated here with flow cytometry using the COPAS Biosort instrument.

### A novel vital staining protocol with BODIPY 493/503 enables the labeling of lipid droplets in vital worms

In addition to fixative staining of worms with BODIPY 493/503, we developed a new vital staining protocol. In contrast to feeding of worms at NGM plates with fluorescent dyes over a long time period, worms were stained in M9/BODIPY 493/503 solution for 20 min. For visualization of BODIPY 493/503 fluorescence in vitally stained worms, microscopy settings were used that did not allow the detection of autofluorescence signals in the green emission spectrum in unstained worms. Confocal three-dimensional (3D) projection images (Fig. 9) demonstrate that vesicles in intestine and hypodermal cells in the head region, the main body hypodermal syncytium hyp7, and the tail region, as well as in gonads and oocytes but not in embryos are stained. To verify that the stained compartments are lipid droplets and not LROs, we performed co-localization studies. First, we determined whether the vital BODIPY 493/503 staining before fixation is in accordance with LipidTox deep red staining after fixation. As shown in Fig. 10A, the vitally BODIPY 493/503-stained structures in head and proximal intestine colocalize with the post-fixation LipidTox-stained lipid droplets. Second, we show that the vitally stained organelles do not overlap with LysoTracker red, a marker for acidic autofluorescent LROs (13, 38) (Fig. 10B). Third, the BODIPY 493/503 signals are in agreement with the CARS signals and do not colocalize with the autofluorescent signals of the LROs (Fig. 10C, and supplementary Fig. IIA). Of note, BODIPY 493/503-feeding worms on NGM plates for 18 h causes dye accumulation in the autofluorescent LROs (supplementary Fig. IIB). Thus, our data indicate convincingly that BODIPY 493/503 labels lipid droplets in vital worms when applied to the worm for a short time period. In Fig. 10D the flow-cytometric analysis of BODIPY 493/503 fluorescence and extinction of vitally stained wild-type N2 and eat-2, daf-7 and glp-1 (25°C)
mutant worms is shown. The resulting fluorescence distribution patterns of the strains are very similar to those obtained by our fixative BODIPY 493/503 procedure (Fig. 8).

**DISCUSSION**

We characterized the body composition of *C. elegans* fat storage mutants by using novel BODIPY 493/503-based microscopy and flow cytometry protocols. Comparison of images obtained with our protocol and label-free imaging in vivo using CARS microscopy revealed similar intestinal and hypodermal fat distributions and lipid droplet morphology of the N2 worms. The size of the lipid vesicles (<3 µm in diameter) stained with BODIPY 493/503 fits well with that recently reported (39). Colocalization of BODIPY 493/503-stained structures with the label-free CARS and Raman lipid signals also proves that BODIPY 493/503 exclusively stains lipid droplets. This finding is in agreement with the observation of Zhang et al. (14), who found that fixative Nile red shows only the lipid droplet-specific emission spectrum, in contrast to that of the vital staining method with Nile red (14). Fixative Oil Red O and Nile red staining were established as a suitable and quantitative method to stain the main fat stores in *C. elegans* (10, 15). In comparison to the Oil Red O procedure, our BODIPY 493/503 protocol easily detects small lipid droplets because of high sensitivity and resolution of the fluorophore. Particularly in regions with high lipid droplet density, visualization of single droplets is feasible with BODIPY 493/503 stain. Furthermore, the BODIPY 493/503 staining procedure preserves the morphology of lipid droplets because the protocol omits any dehydration steps using isopropanol or other alcohols. It has been reported that isopropanol leads to fusion of small adjacent lipid droplets (40). Thus, the BODIPY 493/503 staining procedure seems to be a reliable method to explore lipid droplet
Fat stores of *C. elegans* morphology in *C. elegans* and may be useful for colocalization studies using immunofluorescence or mCherry reporter.

We developed an easy protocol for vital staining with BODIPY 493/503 that enables the labeling of lipid droplets in vital worms. Common vital fat staining procedures in *C. elegans* showed that the dyes do not colocalize with fixative LipidTox (10) but do with LRO markers (10, 11, 13). In contrast, our BODIPY 493/503 signals in vital worms colocalized with fixative LipidTox signals as well as with CARS signals and do not colocalize with the autofluorescent signals of the LRO markers. The feeding of dyes over a longer period, as used in common vital staining protocols, may lead to metabolism of lipid dyes like...
correlate well with those found by Brooks et al. (15). We transformed the data from Brooks et al. to TAG per phospholipids to make them comparable with our data. After this transformation, we found that the eat-2 mutant contains 60% and the daf-7 mutant contains 169% TAG/phospholipid of the wild-type strain, which are consistent with our own data. The glp-1 mutant (25°C) contains 163% TAG/volume of N2. This is lower than the TAG data (297%) reported by O’Rourke et al. (10) but correlates very well with their Oil Red O staining results (147%) and our own BODIPY 493/503 quantification data (155%) for the glp-1 mutant. Because the data obtained by BODIPY 493/503 flow cytometry represent fat content per volume at a single-worm level, this high-throughput fat assessment method will allow an efficient and accurate screening of C. elegans mutants. Such a screening may be confounded by the number of eggs and embryos stored in the uterus of the hermaphrodites. For example, the eat-2 and glp-1 mutants (restricted temperature) have fewer or no eggs in comparison to that of N2 worms, whereas the daf-7 mutants retain more eggs and early-stage embryos. These differences influence triglyceride as well body volume measurement. Thus, we assumed that the resulting fat content per volume is not substantially influenced by different numbers of eggs. To address our hypothesis, the fat and protein content of the eggs and embryos should be determined in future studies.

In contrast to the TAG to volume ratio that seems to be specific for each mutant, protein-to-volume ratios reflecting the proportion of fat-free mass are very similar when comparing different conditions and strains. In support of this, Jonassen et al. (43) showed that wet weight-to-protein ratio (11.8) is constant between larvae and adults. This ratio is in good agreement with our protein-to-volume ratio (12.5), assuming a worm density of ~1 g/cm$^3$. Furthermore,
Fat stores of *C. elegans* 1291 to-protein ratio of 0.2. This is in contrast to that of other species. For example, rats have a TAG-to-protein ratio of 0.8 (48). However, the low fat content and high protein level of *C. elegans* reflects the low fat-to-protein ratio of *E. coli*, the food source of the worms (15). Fatty acid absorption from diet contributes essentially to saturated fatty acids palmitic (C16:0) and sterinic acid (C18:0) content in the nematode, because de novo synthesis is very low (5% and 7%, respectively), whereas the de novo synthesis of MUFAs and PUFAs is up to 20% (42). Brooks et al. (42) showed that there is an inverse correlation of C17iso fatty acid and TAG. C17iso fatty acid is de novo synthesized from the branched chain amino acid leucine. A selective protein deprivation as it is given in *pep-2* mutants that lack intestinal peptide transporter activity (49) results in a decrease in C17iso fatty acid and increased TAG storage, whereas an overall dietary restriction (e.g., in the *eat-2* worm) does not change the C17iso fatty acid concentration but reduces the content of TAGs in total lipid (1) or volume (our data). Also, in mammals, high-protein diets result in a shift to a higher ratio between lean and fat mass (50) and a reduction in food energy intake (51). Hereby, particularly leucine stimulates protein synthesis in muscle (52) and mediates hormonal hypothalamic responses by activating mTOR signaling (53, 54).

Houthoofd et al. (44) found similar protein-to-volume ratios for the N2 wild type and the *eat-2* (ad1115) mutant. Protein is also commonly used as a reference value for TAG. Our data for N2 worms (0.16–0.23 µg of TAG/µg of protein) are supported by other enzymatically determined TAG levels: 0.2 µg of TAG/µg of protein (45) and 0.12 µg of TAG/µg of protein (46). TLC/GC measurements revealed a ratio of 0.15 (7). Brooks et al. reported lower ratios (~0.06–0.07), and Srinivasan et al. reported 10-fold higher values (47). Differences in ratios may be due to different extraction and analysis methods and may be dependent on the developmental stage of nematodes or composition of *E. coli* food source.

Based on the literature and our own data, we estimated the average body composition of adult wild-type N2 worms. The *C. elegans* wild type consists of ~16 ng of TAG/nl, which is about 1.6 w/v % of TAG (w/v %). Assuming a phospholipid fraction of 45%–60% of total lipid, the worm contains 3–4 w/v % lipids. The proportion of protein (~78 ng/nl) is about 7.8 w/v %. Assuming a water content of about 80%–85% (43), there is a difference of 5 to 10 w/v % that may be carbohydrates and insoluble matter, such as components of cuticula. Taken together, the *C. elegans* wild type contains about 5-fold less TAG (1.6 w/v %) than protein (7.8 w/v %), resulting in a TAG-to-protein ratio of 0.2. This is in contrast to that of other species. For example, rats have a TAG-to-protein ratio of 0.8 (48). However, the low fat content and high protein level of *C. elegans* reflects the low fat-to-protein ratio of *E. coli*, the food source of the worms (15). Fatty acid absorption from diet contributes essentially to saturated fatty acids palmitic (C16:0) and sterinic acid (C18:0) content in the nematode, because de novo synthesis is very low (5% and 7%, respectively), whereas the de novo synthesis of MUFAs and PUFAs is up to 20% (42). Brooks et al. (42) showed that there is an inverse correlation of C17iso fatty acid and TAG. C17iso fatty acid is de novo synthesized from the branched chain amino acid leucine. A selective protein deprivation as it is given in *pep-2* mutants that lack intestinal peptide transporter activity (49) results in a decrease in C17iso fatty acid and increased TAG storage, whereas an overall dietary restriction (e.g., in the *eat-2* worm) does not change the C17iso fatty acid concentration but reduces the content of TAGs in total lipid (1) or volume (our data). Also, in mammals, high-protein diets result in a shift to a higher ratio between lean and fat mass (50) and a reduction in food energy intake (51). Hereby, particularly leucine stimulates protein synthesis in muscle (52) and mediates hormonal hypothalamic responses by activating mTOR signaling (53, 54).
free ratio is highly variable depending on macronutrient intake (55). An intriguing question is whether the body composition in C. elegans is variable under different dietary protocols such as reduction of protein content in combination with high carbohydrate or high fat diets.

In conclusion, we developed BODIPY 493/503-based approaches in order to characterize fat storage and body composition of C. elegans mutants. The procedures have a number of advantages. First, the worm anatomy and the native morphology of lipid droplets are preserved. Second, the fluorescent dye allows an accurate and very sensitive quantification of body fat. Third, flow cytometry using BODIPY 493/503-stained worms displays fat quantity per volume at a single-worm level, allowing an accurate screening of fat storage mutants. Using these methods, we characterized the body composition of C. elegans and found that the fat content per volume is specific for dietary restriction, TGF-β, and germline mutants. In contrast, the proportion of fat-free mass is constant among these strains. Together, BODIPY 493/503-based and flow cytometry approaches allow sensitive and accurate assessment of adiposity in large C. elegans populations at a single-worm level.

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