Three-dimensional behavioural phenotyping of freely moving C. elegans using quantitative light field microscopy

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Abstract

Behavioural phenotyping of model organisms is widely used to investigate fundamental aspects of organism biology, from the functioning of the nervous system to the effects of genetic mutations, as well as for screening new drug compounds. However, our capacity to observe and quantify the full range and complexity of behavioural responses is limited by the inability of conventional microscopy techniques to capture volumetric image information at sufficient speed. In this article we describe how combining light field microscopy with computational depth estimation provides a new method for fast, quantitative assessment of 3D posture and movement of the model organism Caenorhabditis elegans (C. elegans). We apply this technique to compare the behaviour of cuticle collagen mutants, finding significant differences in 3D posture and locomotion. We demonstrate the ability of quantitative light field microscopy to provide new fundamental insights into C. elegans locomotion by analysing the 3D postural modes of a freely swimming worm. Finally, we consider relative merits of the method and its broader application for phenotypic imaging of other organisms and for other volumetric bioimaging applications.

Introduction

Since it was proposed as a model organism several decades ago [1] the model organism C. elegans has become an important research tool used in many different fields from behavioural genomics [2] and neuroscience [3] to drug screening [4]. In particular, it is widely used in imaging-based behavioural studies in which the phenotypic effects of genetic mutations and external stimuli are assessed through variations in worm posture and locomotion [5]. Limitations in current imaging techniques mean that these experiments are typically restricted to two dimensional (2D) visualisation and analysis of worms crawling on the surface of an agar gel. However, even on a flat surface, to fully characterise aspects of behavioural response, such as
head movement, requires visualisation and analysis of the shape and motion of the organism in three dimensions (3D). Further, the artificial setting of an agar plate fails to reproduce the complexity of the 3D environment inhabited by *C. elegans* in nature [6], limiting the freedom of the organism to express its full range of behaviours. Real time imaging of the organism in 3D offers the possibility of removing these constraints, allowing us to observe and quantify a greater range of behavioural phenotypes.

Despite significant developments in optical microscopy techniques in recent years [7], fast, minimally invasive volumetric imaging of biological systems remains challenging. Most conventional rapid 3D imaging techniques, such as light sheet microscopy [8] and spinning disk confocal microscopy [9] are inherently sequential, with an image volume built up from a series of image planes captured at different times. This fundamentally limits their ability to capture very fast dynamic processes, such as the unconstrained movement of *C. elegans*. Most of these methods also require the use of fluorescent labels to visualise the structure of interest. An alternative approach to fast 3D imaging is to capture multiple views of the sample simultaneously. In multifocal microscopy this is achieved using a curved diffraction grating which allows multiple focal planes to be imaged onto the camera simultaneously [10], however the number and separation of captured image planes is limited and correcting dispersion effects associated with broadband light sources can be challenging. Capturing images along two [11] or three [12] orthogonal view directions using multiple cameras has been shown to be effective for visualising the 3D posture and motion of *C. elegans*. However, such an approach requires a specialised optical set up with multiple synchronised cameras, along with associated alignment, calibration and image reconstruction processes. The need to illuminate and view the specimen clearly from multiple perpendicular view directions also imposes constraints on sample mounting and the experimental geometry.

Light field microscopy (LFM) [13] is an alternative volumetric imaging method, in which multiple perspective views of the sample are captured simultaneously through a single microscope objective lens. This is achieved using a microlens array (MLA) mounted at the native image (camera) plane, making it both simple and inexpensive to implement on a conventional widefield optical microscope system. Each raw camera exposure (light field image) contains both spatial and angular information, which can be manipulated to generate different perspective and focused views [14] of the object. LFM is suitable for viewing specimens under a range of imaging modalities including bright field, dark field and fluorescence [15]. In this article we describe how combining LFM with computational depth estimation methods, originally developed for photography [16], allows measurement of the 3D motion and body shape of *C. elegans* at a speed limited only by the frame rate of the camera. We apply this method to analyse 3D behavioural phenotypes by quantifying differences in posture and locomotion between two collagen mutants (*dpy-10* and *dpy-13*) freely moving within a 3D gel. By extending the eigenworm method of postural analysis [17] to 3D we use quantitative LFM to investigate the 3D swimming motion of a wild type worm. Finally, we discuss the broader application of quantitative LFM which, as a flexible imaging technique readily implemented on most standard research microscopes, offers new possibilities for experimental investigation of *C. elegans* and other dynamic 3D biological systems.

**Methods**

**Light field microscope system**

In order to capture 3D body shape and locomotion phenotypes of freely moving *C. elegans* we developed a custom LFM system by modifying a standard upright widefield microscope (BX51WI, Olympus) (**Fig 1**). The properties (lateral and angular resolution, depth of field and field of view) of an LFM system are strongly dependent on the parameters of the microlens...
array and its location in the imaging pathway (see supporting information S1 Text). For experi-
mental and computational simplicity we adopt the ‘classic’ LFM configuration first described by
Levoy et al. [13] by mounting an MLA (MLA-S125-F20, RPC Photonics) in the native image
plane and projecting a 1:1 image of the back focal plane onto the image sensor of a scientific
CMOS camera (ORCA-Flash 4.0v2, Hamamatsu Photonics) using a pair of achromatic doublet
lenses (Thorlabs Inc.). Using a 10x/0.3 objective lens (UMPLFLN, Olympus) this gives a field of
view of 1.3 mm x 1.3 mm. A field lens (a plano-concave singlet with a focal length of half the mag-
nitude of the positive doublets) mounted immediately in front of the camera sensor was used to
improve imaging of off axis image points by reducing Petzval field curvature. All these auxiliary
components were mounted on an optical breadboard, supported on the same optical table as the
microscope system. In this configuration each light field image (Fig 1—inset) is comprised of a
tiled array of circular microlens subimages. Selecting the same pixel from each subimage creates a
‘pinhole’ view of the object from a corresponding view direction, whereas summing over all the
pixels within each subimage results in an image focused at the native image plane. A captured
light field $L_o$ can be computationally refocussed, to simulate the effect of an axial shift in sample
position, by shearing it parallel to the $(x, y)$ plane using $L_o(x, y, u, v) = L_o(u(1 - 1/\alpha) + x/\alpha, v(1 - 1/}$
$a) + y(a,u,v)$ [14]. This procedure, equivalent to laterally shifting each pinhole view prior to summation, allows computation of a focal series ($z$-stack) of images from a single camera exposure.

To increase image contrast and improve the reliability of image segmentation the microscope was configured for differential interference contrast imaging, with matched pairs of polarisers and Normarski prisms in the optical path before and after the sample. All images were captured at 100 frames per second (FPS), the maximum full frame rate of the camera, to minimise motion blur. For 3D reconstruction and behavioural analysis time lapse image sequences were down sampled to 20 FPS and exported as 16 bit multipage tiffs.

**Image reconstruction and depth estimation**

All images were reconstructed and analysed using custom software written in MATLAB (MathWorks). Images were first corrected for vignetting, which decreases the detected radiation for off axis rays, by dividing each raw light field by a corresponding background image captured under identical illumination and detection conditions. The 2D light field image was then reshaped into a four-dimensional array, $L_0(x,y,u,v)$ using an automatic rectification procedure in which an initial estimate of the pitch and orientation of the subimages was from obtained by detecting the prominent peaks in a 2D Fourier transform of the raw image. This estimate was used to create a corresponding binary mask of tiled subimages. The mask parameters (pitch, orientation, lateral offset from centre), were then fine-tuned by translating and scaling the mask in to maximise the overlap (2D cross correlation) with the original normalised image light field image. The final mask settings were then used to resample the light field with an integer number of pixels in each subimage and reshape it into a 4D array.

Variation in image features with (post capture) changes in perspective and focus provide two ways of visualising the axial ($z$) position of objects within a light field image. The direction and magnitude of the apparent lateral shift in an object’s position with view direction (parallax) indicates how far it lies from the native object plane. Similarly, changes in sharpness of an image feature with computational refocusing indicate the position of the corresponding object relative to the native object plane. By analysing how these attributes of the light field change as it is computationally refocused (sheared parallel to the ($x$, $y$) plane) it is possible to determine the three-dimensional position of features within the sample. Importantly, the applied shear parameter, $\alpha$, is related to the axial position in the sample by

$$\Delta z = P(1 - 1/\alpha)F/M^2,$$

where $P$ is the subimage pitch in camera pixels, $F$ is the focal length of the MLA and $M$ is the transverse magnification of the microscope.

In order to reconstruct the 3D body shape of the worm in each image frame we applied a depth estimation method originally developed for light field cameras [16], in which depth estimates based on defocus and correspondence are computed separately and then combined to produce a single (more robust) estimate. In practice this amounts to refocusing the light field and then computing defocus and correspondence responses as a function of the shear parameter $\alpha$ (see supporting information S1 Fig). A defocus depth response value is computed for each pixel in a refocused image, $L_s(x,y) = \sum_{u,v} L_s(x,y,u,v)$ over a window $W_D$ using, $D_s(x,y) = 1/|W_D| \sum_{x,y \in W_D} |\Delta_{xy} L_s(x,y)|$, where $\Delta_{xy}$ is the Laplacian operator. The correspondence depth response is defined as the average angular deviation in pixel value over a window $W_C G_s(x,y) = \frac{1}{D_s(x,y)} \sum_{x,y \in W_C} \sigma^2_{xy}$, where $\sigma^2_{xy} = \frac{1}{D_s(x,y)} \sum_{x,y \in W_C} (L_s(x,y,u,v) - L_s(x,y))^2$. In both cases we found that a window size of five pixels gave optimal results. Defocus and correspondence depth estimates, $\alpha_D$ and $\alpha_C$, are defined as the maximum and minimum of the defocus and
correspondence responses, \( z_p^+(x, y) = \arg \max_y D_p(x, y) \) and \( z_p^-(x, y) = \arg \min_y C_p(x, y) \). The final depth value is then computed by combining the depth estimates and corresponding confidence values, defined as the ratio of the defocus and correspondence response at the optimal depth to the corresponding value at the next best depth (next largest/smallest local maximum/minimum), using a Markov Random Fields optimisation [16].

An estimate of the 3D body shape of the organism in a given image frame was computed by combining a depth image with a corresponding segmentation mask which defines the lateral footprint of the organism (projection along the z axis) (see Fig 2 and supporting information S1 Video). An approximate binary segmentation mask was generated from the on axis pinhole view by adaptive intensity thresholding, morphological hole filling and closing. This mask was then smoothed using active contours to create a final segmentation mask (Fig 2(A)). To improve the depth estimate a two frame temporal Kalman filter was applied to remove spurious depth values. A midline skeleton was then defined by fitting a smoothing spline to the 3D coordinates of pixels lying along a line from the nose to the tip of the tail through the centre of the mask (Fig 2(C)).

We found this method was effective for generating mid-line skeletons except when the worm adopted a self-occluded posture by coiling back across itself (see supporting information S2 Video). In self-occluded regions it was no longer possible to assign a single depth value to each lateral image pixel and both depth cues returned ambiguous results. For this reason, each time-lapse light field dataset was visually assessed prior to depth estimation and those containing self-occluded postures were excluded from further analysis.

By far the most computationally intensive part of the analysis was in computing the depth estimate for each image feature. The time taken for this step depended on a number of factors including the axial range over the depth responses are computed. On a desktop PC with an Intel Xeon E5-2690 v3 12 core 2.6 GHz processor and 64 GB of RAM, it took 24 seconds per time point to estimate depth of features over 1 mm, or approximately 8 minutes per second of video analysed at 20 frames per second. We anticipate that significant speed improvements should be possible by optimising software for computational efficiency. Further, the process is highly amenable to parallel processing and we anticipate that it could be speeded up considerably by exploiting the computing power of a graphics processor unit.

Analysis of 3D posture and motion
To quantify posture and locomotion the metrics shown in Table 1 were computed directly from the 3D mid-line skeleton of the worm in each image frame. To compare behavioural phenotypes of dpy mutants these values were averaged over all time lapse image sequences.
Calculation of eigenworms and analysis of postural modes

Eigenworms were computed following the method described in [17]. Midline body skeletons were split into 25 segments, with the orientation of each segment described by a pair of angles \((\theta, \phi)\) (Fig 2(C)). The mean of each of these angle vectors over the body length, \(h_\theta\) and \(h_\phi\), was set to zero, giving a shape representation which depends only on the posture adopted by the worm and rotation about its long (roll) axis. All 25 element postural angle vectors successfully extracted from midline skeletons for wild type, \(dpy-10\) and \(dpy-13\) worms moving in agarose and swimming wild type worms were concatenated, before eigenworms were computed using the MATLAB function ‘pca’. In practice, the close similarity between eigenworms computed from azimuthal and polar angle vectors allowed use of a single common set of eigenworms, which we chose as the azimuthal set. Eigenvalues for swimming motion were determined for each light field image frame by projecting the extracted midline skeleton onto this common eigenworm basis.

In contrast to alternative ways of describing 3D posture, such as the Frenet-Serret equations, we found this simple approach was relatively insensitive to noise in the midline skeleton and gave a robust and intuitive description of body shape. Computing a single set of eigenworms by combining azimuthal and polar angle vectors could in principle give a more efficient representation of the 3D posture, however the resulting projections are not simple to interpret and there remain questions about how best to normalise the angle vectors. As well as allowing a direct comparison with 2D postural mode analysis, treating the azimuthal and polar angles independently allows an independent validation of depth estimation by comparing corresponding azimuthal and polar eigenworms. Computing separate lateral and axial postural modes may also be important for particular behavioural assays [18].

### C. elegans culture and preparation

\(C.elegans\) strains, N2 (wild-type), CB128 (\(dpy-10\)) and CB184 (\(dpy-13\)), were obtained from the Caenorhabditis Genetics Center (University of Minnesota) and maintained according to standard procedure [1]. For experimental purposes, L4 staged animals were transferred to fresh plates a day prior to imaging (by which time they had reached adult stage). To visualise organisms moving in 3D, worms were prepared in an agarose gel (at 0.25% wt/vol in M9 buffer solution [19]) on a microscope coverslip.

To calibrate the depth scaling factor a single worm was paralysed by treatment with BDM (2,3-butanedione monoxime), diluted to 15 mg/ml in M9 buffer, for 60 minutes. For visualising swimming behaviour several drops of M9 buffer solution were added to an agarose pad containing a number of wild-type adults, causing worms to detach from the surface. A time

### Table 1. Posture and locomotion metrics computed to compare behaviour of \(dpy-10\) and \(dpy-13\) mutants.

| Metric                                      | Formula                                                                 |
|---------------------------------------------|------------------------------------------------------------------------|
| Centroid speed \(CS_i\)                    | \(CS_i = |r_{i+1} - r_i|/(t_{i+1} - t_i)\), where \(r_i\) is the centroid position vector at time \(t_i\). |
| Non-planar deviation (NPD)                 | \(NPD = R_3/R_1\), where \(R_1\) and \(R_3\) are the lengths of the longest and shortest principal axes of the best fit ellipsoid to the worm posture, computed by performing principal component analysis on the corresponding mid-line skeleton. |
| Curving rate for forward runs \(CR_i\)     | \(CR_i = \frac{1}{|r_{i+1} - r_i|} \cos^{-1} \frac{(|r_{i+1} - r_i|)}{R_1(R_3)}, \) where \(r_i\) is the centroid position vector at time \(t_i\). |
| Directional autocorrelation for forward runs | \(D_n = \frac{1}{N} \sum_{i=1}^{N-1} v_i \cdot v_{i+n}\), where \(v_i\) is the unit direction vector in the \(i\)th frame and \(N\) is the total number of images in the sequence. |

[10.1371/journal.pone.0200108.t001](https://doi.org/10.1371/journal.pone.0200108.t001)
lapse light field sequence of duration 30 seconds was captured beginning as soon as an animal was positioned close to the centre of the field of view of the microscope.

Results

Calibration of depth scaling factor

The depth estimation accuracy was tested using a series of light field images of a paralysed *C. elegans* specimen mounted on a microscope coverslip captured as the microscope focusing stage was displaced by known amounts. Fig 3 shows the estimated depth (averaged over the midline of the worm) versus known focus offset (measured depth), where defocus and correspondence estimates have been offset by +/- 100 μm for clarity. Both correspondence, defocus and combined depth estimates vary linearly with real depth ($R^2 \geq 0.9989$ for all three) over the entire measured range of +/- 1 mm. The gradients of individual depth estimates are less than one, which we attribute to differences between the nominal and real system parameters used to relate the light field shearing parameter ($\alpha$) to depth. All subsequent final depth estimates were scaled by the gradient of the fit to the combined depth estimate (0.883). The root mean square error of the linear fit to the combined estimate is 11.1 μm, which is a measure of the depth sensitivity (axial resolution) of the system. This value is considerably smaller than both the DOF of the refocused images (65 μm) and the total thickness of the worm body (approximately 50–70 μm for an adult), thus the depth value assigned to a given part of the worm corresponds to a contrast weighted average over the total worm thickness.

![Fig 3. Calibration of depth scaling factor. Correspondence (red), defocus (blue) and combined (black) depth estimates versus known depth obtained by capturing a series of light field images as a paralysed *C. elegans* specimen was translated through a series of known axial positions. Defocus and correspondence estimates offset by + 100 μm and -100 μm for clarity.](https://doi.org/10.1371/journal.pone.0200108.g003)
Real time volumetric imaging of *C. elegans*

For an initial proof of principle test, we captured a 30 second long time lapse light field image sequence of an adult wild type worm moving inside a block of agarose gel. During this period the worm underwent four phases of forward locomotion to reach different locations within the gel. At the end of each forward phase the worm paused briefly and executed a series of exploratory head movements (foraging) before reversing. Fig 4(A) shows on axis pinhole views of the worm in three second intervals. Fig 4(B) shows all 600 mid-line skeletons extracted from the light field images captured during this period (nose tip indicated in red). The grey bounding box shown in the isometric projection in the upper left part of Fig 4(B) indicates the total volume of 3D space visited by the worm during the sequence (0.027 mm$^3$). The xy and xz projection views clearly illustrate the 3D movement of the head during the foraging phase, with the nose exploring a volume of 0.009 mm$^3$ during the sequence. Such 3D information is a useful addition to existing methods for detecting and analysing foraging behaviour [20].

Quantitative comparison of 3D behavioural phenotypes of cuticle mutants

As a proof of principle experiment we applied quantitative light field microscopy to analyse the behaviour of two *C. elegans* cuticle mutants, *dpy-10* and *dpy-13*. We have previously studied these two mutants and found significant differences in cuticle topography [21]. Both have mutations in genes encoding collagen proteins, similar gross morphologies and are shorter...
and fatter than wild-type worms. However, whilst the cuticle of dpy-13 contains regular annuli and furrows, dpy-10 lacks this organised structure (see Fig 5). The cuticle of C. elegans has an important role in the animal’s ability to move in complex environments such as soil [22, 23]. The functional role of the annuli are not known, but are thought to permit flexibility during movement [22, 24]. In order to elucidate whether the absence of the annuli structure alters posture and locomotion in 3D environment, we compared dpy-10 and dpy-13 worms moving within agarose gel.

For each data set imaging was begun immediately after the application of a gentle mechanical stimulus (tap) to the sample mount to stimulate movement of the worm. Time lapse light field images were then acquired until the worm moved outside the field of view of the microscope. In total we recorded and analysed 27 separate time lapse image sequences of seven different dpy-10 animals with a total duration of 9.2 minutes and 19 image sequences of 10 different dpy-13 animals with a total duration of 4.4 minutes.

To quantify the posture and locomotion of the worms in each image sequence we computed the centroid speed, non-planar deviation (NPD) and curving rate in each image frame and the directional autocorrelation during forward runs (for details see Table 1). NPD is a measure of how far the worm’s posture extends away from a best fit plane and quantifies the ability of the worm to bend simultaneously in two perpendicular directions, whilst curving rate and directional autocorrelation describe how rapidly the worm changes its direction of motion. The averaged results (Fig 6) indicate that both mutants move with a similar speed, with mean centroid speeds of 46.1 ± 5.7 μm/sec and 47.0 ± 6.7 μm/sec for dpy-10 and dpy-13 respectively. Both the mean curving rate and NPD are substantially lower for dpy-10, suggesting that the
lack of regular annuli and furrows in the cuticle limit the worm’s ability to rapidly change direction and adopt complex postures and supporting the theory that the cuticle annuli give flexibility to the worm during movement. This is also suggested by the more rapid decay in the directional autocorrelation curve for dpy-13 and exponential fits \( D(t) = Ae^{-bt} \) to the data yield decay constants \( b \) of 0.20 \( \pm \) 0.04 and 0.33 \( \pm \) 0.08, however these differences are not statistically significant \( p = 0.16 \).

3D postural modes of swimming animals

An alternative method to analyse worm behaviours is by describing the associated body shapes in a space of fundamental postures [2]. In this case, a 2D worm posture is represented by a vector of tangent angles describing the orientation of different sections of the mid-line skeleton. Performing principal component analysis on a large set of these tangent vectors, derived from time-lapse images, yields a set of fundamental postural modes or eigenworms [17]. Projecting a given midline skeleton onto this basis allows the posture to be described in terms of the associated projection amplitudes or eigenvalues. By capturing the 3D worm bodyshape quantitative light field microscopy allows this method to be extended to measure the 3D postural modes of freely moving worms. We achieved this by representing each reconstructed 3D midline skeleton as a pair of 25 element vectors \( \theta(s) \) and \( \phi(s) \) containing azimuthal and polar tangent angles. We generated 3D eigenworms from 3D midline skeletons for dpy-10, dpy-13 and wild-type worms in agarose. Fig 7(A) shows the first four eigenworms derived from azimuthal and polar tangent angle vectors. Note the close resemblance between azimuthal and polar eigenworms which may be considered a further validation of the accuracy of the depth estimation method. This close similarity also means that in practice a single set of eigenworms (derived from either the azimuthal or polar tangent vectors) can be used to represent motion in lateral and axial directions. Similarly to previous work based on 2D eigenworms [17], our results indicate that 95% of the body shape variation is captured using the first four eigenworms in \( \theta \) and \( \phi \) (Fig 7(B)).

Analysis of 2D eigenworms [17] has shown that the joint probably density function of the first two eigenworms forms a ring structure, indicating that the first two eigenworms form an oscillator with an approximately fixed amplitude and varying phase describing the basic crawling behaviour of the animal. We observe a similar result in 3D, however allowing the animal to move in 3D means that amplitude of the oscillator varies with time (depending on the orientation of the animal) and the resulting trajectory in the \((a_1, a_2)\) plane is elliptical. Fig 8(A) shows

![Fig 7. 3D eigenworms.](https://doi.org/10.1371/journal.pone.0200108.g007)
examples of these elliptical trajectories for a wild type *C. elegans* swimming in buffer solution. We can gain further insight into the 3D swimming motion of this animal by considering the relationship between the corresponding $\theta$ and $\phi$ projection amplitudes. The top left panel in Fig 8(B) shows the first $\theta$ and $\phi$ eigenvalues for the same swimming wild type animal. The data bear a striking resemblance to the rose or rhodonea curve, $r = A \cos(kt + c)$, suggesting that we can write:

$$a_\theta = A \cos[k(at + c)] \cos[at + c]$$

$$a_\phi = A \cos[k(at + c)] \sin[at + c]$$

Performing least squares fits to Eq 2 yields a good match to experimental data (Fig 7(C) and 7(D)), indicating that this simple expression effectively describes the 3D swimming motion of the organism. If the first cosine term in each expression is associated with the ventral-dorsal oscillatory motion of the worm body during swimming, then the second term describes the frequency with which the animal changes the orientation of this plane of oscillation. Independent fitting to the experimental data for the first and second eigenworms yields a mean value for $k$ of 8.5 with a standard deviation of 0.3. To our knowledge this is the first time that this ‘rolling frequency’ has been experimentally measured.
Discussion

We have shown that combining LFM with computational depth estimation allows quantification of 3D postural and locomotion characteristics in *C. elegans*, permitting extended phenotypic imaging of the organism in a more natural setting. In contrast to other 3D microscopy techniques, light field microscopy is readily implemented on many of the wide-field microscope systems currently in use using widely available and relatively inexpensive optical components. The relative simplicity and flexibility of the technique make it attractive for quantitative 3D imaging of a range of other dynamic biological systems including other model organisms, such as the *Drosophila* embryo, and 3D cell cultures and organoids. The capability of quantitative LFM to capture 3D sample morphology from label free images makes it particularly attractive for long duration time-lapse imaging of sensitive biological samples for which the use of fluorescent labels and high intensity laser excitation induces undesirable phototoxic effects [25]. However, LFM is also suitable fast volumetric fluorescence imaging and has been used for neural activity imaging in *C. elegans* and other organisms using fluorescent calcium indicators [26, 27]. This potential to capture 3D behavioural and neural activity information simultaneously opens up exciting new possibilities in experimental neurobiology. The primary limitation of LFM is its relatively modest spatial resolution, which stems from the need to sacrifice lateral spatial information in order to simultaneously capture multiple perspective views, and this has limited the application of the technique. There are, however a number of strategies for increasing spatial resolution. Firstly, changing the properties of the microlens array allows a trade-off between the spatial and angular sampling rates and the depth over which post capture refocusing is possible. Actively maintaining the object within the field of view using a closed loop object tracking system [28] would allow longer duration time-lapse imaging, permitting a reduced field of view and the use of higher magnification objective lenses for improved spatial sampling and achievable resolution. Exploiting the higher spatial sampling rate away from the native image plane by reconstructing light field images using 3D deconvolution [29], offers another way to improve spatial resolution. However, in preliminary experiments using this latter approach we found that the strong dependence of the spatial resolution on object depth and the presence of deconvolution artefacts caused errors in depth estimation. With further development of the depth estimation algorithms it may be possible to overcome these limitations. Recent work [30, 31] has shown mounting the microlens array in the pupil plane of the microscope objective rather than the image plane can improve lateral spatial resolution. Given the limited number of camera pixels this typically comes at the expense of fewer angular samples, but it would be an interesting exercise to compare the ability of these different LFM configurations to capture and reconstruct 3D samples structures.

A further technical limitation of LFM stems from the relatively limited angular baseline, which makes depth estimation and 3D reconstruction of complex, self-occluded postures, such as coils and knots, difficult. Recent work [32] has shown that matching a 2D image to a superposition of eigenworm postures offers a way to track coiled body shapes and such an approach may prove useful for reconstructing some self-occluded 3D body shapes.

Supporting information

S1 Text. Spatial resolution and working volume in LFM. A discussion of how microscope design parameters affect the lateral spatial resolution, field of view and effective depth of field in a light field microscope.

(DOCX)
S1 Fig. Schematic illustration of computational depth estimation process. The central part of the figure depicts the processing of light field shearing in 3D (equivalent to laterally shifting each pinhole view). The left panel shows the refocused images created by integrating the sheared light field over all view directions (u, v). The right panel shows three 2D epipolar images, with each row of pixels corresponding to a line profile through a single pinhole image. When the light field is sheared by the amount corresponding to the depth of the object, its position does not change with view angle and it appears as a vertical line in the epipolar image. In this example defocus and correspondence responses (blue and red curves), both indicate the worm body is offset by ~-200 μm from the native object plane.

(TIF)

S1 Video. Lateral segmentation and depth map for a freely swimming wild-type worm. Left: on axis pinhole view (grayscale) and segmented outline (yellow) of a wild-type worm swimming in buffer solution. Right: corresponding combined depth map. Images displayed at 20 frames per second.

(MP4)

S2 Video. Examples of self-occluded postures adopted by C. elegans. Each image sequence shows the on axis pinhole views computed from part of a longer duration light field time lapse image set of a wild-type worm moving in agarose gel. In self-occluded (overlapping) regions it is no longer possible to assign a single depth value to each lateral image pixel and both depth cues return ambiguous results preventing reliable 3D reconstruction.

(MP4)

Acknowledgments
This work was funded by Advanced Grant 247401 from the European Research Council (ERC-2009-AdG, MicroNanoTeleHaptics), and a Capital grant for Great Technologies grant (EP/K005030/1) from the UK’s Engineering and Physical Sciences Research Council. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

The authors thank the Caenorhabditis Genetics Center (CGC), which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440), for providing dpy mutant C. elegans strains.

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