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High temporal resolution measurements of movement reveal novel early-life physiological decline in C. elegans

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Abstract

Age-related physiological changes are most notable and best-studied late in life, while the nature of aging in early- or middle-aged individuals has not been explored as thoroughly. In C. elegans, many studies of movement vs. age generally focus on three distinct phases: sustained, youthful movement; onset of rapidly progressing impairment; and gross immobility. We investigated whether this first period of early-life adult movement is a sustained “healthy” level of high function followed by a discrete “movement catastrophe”—or whether there are early-life changes in movement that precede future physiological declines. To determine how movement varies during early adult life, we followed isolated individuals throughout life with a previously unachieved combination of duration and temporal resolution. By tracking individuals across the first six days of adulthood, we observed declines in movement starting as early as the first two days of adulthood, as well as high interindividual variability in total daily movement. These findings suggest that movement is a highly dynamic behavior early in life, and that factors driving movement decline may begin acting as early as the first day of adulthood. Using simulation studies based on acquired data, we suggest that too-infrequent sampling in common movement assays limits observation of early-adult changes in motility, and we propose feasible strategies and a framework for designing assays with increased sensitivity for early movement declines.

Introduction

Of many physiological changes that occurs during aging, loss of mobility is a critical factor in loss of health and well-being. Changes in mobility in humans are associated with changes in other functional domains (e.g. cognition [1, 2]), and decreased mobility predicts future mortality in both humans [3, 4] and model organisms [5, 6]. Given the importance of mobility and the ease of measuring mobility during aging in humans and model organisms, considerable effort has gone into the study of mobility throughout life.
Similar to humans, the model organism *C. elegans* shows stereotypic age-related changes in movement and neuromuscular function. This makes it a very tractable model with which to study the timecourse of age-related changes in mobility. Physiological processes seen in humans have also been implicated in movement decline of *C. elegans*, such as sarcopenia [5, 7] and age-related deterioration of neurons [8–10].

While such patterns of movement decline are well-characterized in later adult life, comparatively little work has investigated changes in movement function early in adulthood. This is surprising given that early adulthood is a time of great change in many aspects of animal physiology, including: continued changes in size; transition into and out of reproductive maturity [11]; changes in food consumption [12] and defecation [13]; and increased propensity for extrinsic damage from environmental factors such as colonization by bacterial food [14]. Previous studies of age-related changes in *C. elegans* movement often examine movement decline as a progression of stereotyped behavioral classes: a period of consistently high, vigorous activity in early life (approximately days 1–6 of adulthood at 20˚C); followed by rapid decline and loss of spontaneous movement; and eventually gross immobility even to external stimulus [5, 15, 16]. These qualitative classes have been borne out to varying extents in quantitative studies of movement behavior. Some studies have broadly recaptured these observations (i.e. an early life maintenance of relatively constant, high physiological function) [17, 18], while others suggest that movement decline may begin earlier in adulthood [19–21] (though the onset of this decline varies between studies).

One plausible reason why early-life movement appears stable in some studies is that movement assays are not optimized to discern between different levels of vigorous activity. On standard vermiculture plates, animals can travel at high speeds (up to ~500 μm/s) [22], making observation of movement challenging. A number of strategies/technologies have emerged to track animals, ranging from video imaging with custom tools for tracking animals on plates [23, 24] to novel culture systems with specialized imaging hardware [25, 26]. In many analyses, especially longitudinal studies using custom systems/software, investigators typically pursue one of two strategies: obtain frequent, low-quality measurements of movement (a handful of images obtained every few hours) [21]; or obtain high-quality movement measurements relatively infrequently (e.g. once-daily videos of several minutes) [17, 19, 25, 27]. The latter strategy has historically been more common. It is not clear, however, the degree to which movement measures from either of these two strategies correlate with “perfect” (if unobtainable) observations of movement: continuous, lifelong video-recording of every individual. As such, it is also not clear in what circumstances a failure to observe changes in early-life movement in *C. elegans* would reflect genuine aging biology vs. technical limitations.

We thus investigated whether young-adult animals experience age-related declines in movement by collecting movement data at previously unattained spatial and temporal resolution. To measure longitudinal changes in animal movement at high temporal resolution, we employed the *C. elegans* Worm Corral longitudinal culture system [21, 28]. This system has the salient limitation that each individual is confined to a relatively small area (a circular pad of bacterial food ~1.5–2 mm in diameter). However, this allows us to efficiently make repeated measurement of changes in behavior for multiple, isolated individuals across multiple days. Using this system, we demonstrate that second-by-second measures of movement are surprisingly well-correlated with measurements of animal movement at considerably lower temporal resolutions, even as low as one frame per minute (fpm). We then made minute-to-minute measurements of movement for a cohort of individuals over the course of several days of early adult life. From these data, we found that many, though not all, individuals do exhibit substantial decline in movement during this period. Finally, using simulations based on the minute-
to-minute data acquired in this study, we find that typical protocols used to assess animal movement in standard vermiculture may be insensitive to early-life physiological changes. To address this concern, we propose several simple improvements to increase sensitivity to age-related movement changes. Overall, this study not only suggests the presence of important age-related physiological changes in mobility in early adulthood, but also provides a schema by which investigators can verify the fidelity of data-acquisition strategies and their sensitivity to early-life mobility changes.

Methods

Animal husbandry

For reasons unrelated to this analysis, all experiments in this study use the *C. elegans* strain BA671 *[spe-9(hc88)]*. This strain has a temperature-sensitive fertility defect above 25˚C, but is otherwise phenotypically wild type at 25˚C and completely phenotypically wild-type at 20˚C [29]. Strains were maintained at 20˚C on standard NGM plates seeded with *E. coli* OP50 [30].

Worm corral bacterial food & animal preparation

We maintained our lifelong culture system at 20˚C for these experiments to ensure the movement results would be as comparable as possible to standard assays which are also typically performed at 20˚C. As *spe-9(hc88)* individuals are fertile at this temperature, we prevented progeny accumulation by instead employing feeding RNAi using the RNAi-compatible OP50-derivative strain OP50(*xu363*) [31] transformed to produce dsRNA against the embryonic transcription factor *pos-1*. Animals fed *xu363/pos-1* during late development and early adult life produce fertilized, but developmentally arrested eggs.

Concentrated *xu363/pos-1* bacterial food for Worm Corrals was prepared as follows. First, a culture was prepared by inoculating LB with a single colony of *xu363/pos-1* and incubated at 37˚C for 17 hours (“overnight”). This overnight culture was diluted down to an optical density (OD) of 0.25, grown up to an OD of 0.5, and induced with addition of isopropyl beta-d-1-thiogalactopyranoside (IPTG) at 1 mM final concentration followed by further incubation for 3 hours. The bacterial culture was resuspended in M9 salts solution at ~75% wt/vol and stored at 4˚C for no more than 3 days.

Embryos were prepared for use in Worm Corrals by setting up P0 animals for a synchronized egg-lay. P0s were synchronized at the L4 stage and, the following day, were transferred to a new plate to lay eggs. To prevent cross-contamination between *xu363/pos-1* and OP50 (which doubles faster and would thus otherwise outgrow the *xu363/pos-1*), the egg-lay plate was seeded with a control *xu363* strain transformed to produce dsRNA against GFP, and transferred animals were deposited off the *xu363/GFP* lawn. Embryos were picked from the egg-lay plate 17–18 hours after transfer of P0s. For each experiment, embryos were picked from a single plate to the worm corral device at room temperature, and were out no more than 1–1.5 hours prior to initiation of experiment.

Construction of longitudinal culturing devices

For longitudinal observation experiments, custom solid media longitudinal culturing devices (“Worm Corrals”) were used [21, 28]. Briefly, the solid media in these devices consists of a PEG-based hydrogel containing a modified formulation of NGM (pH 6.3 with 20 μg/mL cholesterol in solution and lacking calcium chloride). The mixed solution was pipetted in a liquid state into a reservoir formed from a 2.5 mm thick aluminum frame affixed to a standard microscope glass slide, and left to cross-link for 1.5 hrs at room temperature. After
crosslinking. 0.5 μl droplets of concentrated xu363/pos-1 bacteria were pipetted onto the hydrogel surface, and a single embryo was transferred from an egg-lay plate into the drop manually with an eyelash to minimize disturbance of the food pad. After loading the hydrogel with bacteria & embryos, the device was sealed by adding a thin layer of polydimethylsiloxane (10:1 base:cure ratio) onto the hydrogel surface.

Devices were placed on the stage of an automated compound microscope during subsequent culturing and observation. The stage was housed in a sealed enclosure that was kept at constant temperature (20˚C) and humidity (> 90% relative humidity) throughout the duration of the experiment.

Data acquisition
During all experiments, standard brightfield imaging was performed with a compound microscope (5× objective) fitted with a Zyla 5.5 sCMOS camera (sensor size 16.6 x 14 mm; pixel size 6.5 μm; Andor Instruments, Belfast, UK). Prior to data collection, animals were allowed to progress through larval development for 48 hours, such that data acquisition began just prior to the beginning of adulthood for all animals. During this initial period, individuals were followed by taking a single brightfield image every three hours to assess for proper development.

For experiments in which data was collected at 1 frame per second (fps), each individual was continuously imaged in standard brightfield conditions for a total of 20 minutes (allowing for a theoretical maximum of 72 animals to be serially imaged during a 24-hour period). For experiments in which data was collected at 1 frame per minute (fpm), all animals were imaged at each timepoint. In these latter experiments, movement was also stimulated every 3 hours by exposure to a blue LED lamp (Spectra light engine; Lumencor, Beaverton, OR; 470 nm center frequency) for 500 ms. However, because this study is restricted to unstimulated movement, data from the 30 minutes following each stimulation were excluded from further analysis.

Post-experimental image processing, data analysis, and data simulation
The location of each animal in every image was identified using a convolutional neural network, as in previous work [32] (Fig 1A). To assess consistency of bacterial food pad sizes, the bacterial lawn was identified by fitting a two-mode Gaussian mixture model to the image intensity histogram, and taking all pixels corresponding to the lower-intensity mode as associated with the (visually darker) lawn. This procedure was performed for the first five images of that image series, and these binary image masks were combined with a bitwise OR operation to obtain a robust estimate of the entire lawn region (Fig 1A and 1B). These masks were verified by manual inspection. Onset of adulthood for each animal was identified manually by an experimenter and all reported times are specified with respect to the first egg laid, 3–6 hours after the exit from the L4 molt [21]. In this study, movement was measured as the change in position of the centroid of the animal across two or more images.

Linear fits were performed using standard linear regression methods. P-values for Pearson linear regression/correlation coefficients were calculated using the Wald test. We defined a correlation as of biological interest if it reached statistical significance at p < 0.05 (corrected for multiple comparisons by Bonferroni correction as appropriate) and yielded a coefficient of determination (R²) > 0.10. Correlations with lower coefficient of determinations, regardless of p-value, do not capture a great deal of the relevant biological effect and empirically have proven difficult to reproduce.

To produce bootstrap confidence intervals of individual movement decline rates (i.e. as in Fig 3D), each individual’s movement data were resampled as follows. For each 3-hour block, each individual’s minute-to-minute displacements within that block were resampled (with
replacement) and summed to generate a resampled estimate of total movement within that block. Then, overall movement decline rate was calculated as the slope of a linear fit across the resampled movement sums for each 3-hour block. This process was repeated 10,000 times for each individual, and a 95% confidence interval was generated from the estimates for the decline rate. Simulated timecourses of movement were generated by subsampling data acquired at 1 fpm. To sample the “Single Snapshot” and the “Hybrid Movement” protocols, individual images/worm positions were taken at the prescribed sampling interval from the data. To simulate the “Daily Video” protocol (15 minutes of images obtained at 1 fps), we simply used 15 minutes of images obtained at 1 fpm, from the beginning of each 24-hour period. Though these two recording regimes seem very dissimilar, we found from our small-scale 1 fps pilot dataset (Fig 2) that there is a strong correlation between total displacement over 15 minutes at 1 fps and net displacement at (subsampled) 1 fpm imaging (R^2 = 0.81; S1 Fig).

**Results**

**Minute-scale measurements of movement correlate with finer animal movement**

To investigate the movement behavior of early-adult animals, we first set out to determine minimal conditions for reliably measuring movement in young animals over multiple days. Ideally, each animal under study would be continuously imaged throughout the experiment at a frame rate and magnification sufficiently high to capture all motion. For experiments, this is generally impractical, as magnification, frame rate, number of animals studied, and frequency of imaging each individual all generally trade off against one another. These trade-offs led us to consider how well an animal’s movement at the second-by-second scale is represented by measurements taken at longer intervals.

We assessed whether minute-to-minute measurements were sufficient to represent finer movement activity. We acquired 20-minute video recordings at 1 fps of day 1 adult individuals, isolated from one another via the Worm Corral system (Figs 1 & 2A). Based on previous studies, video imaging at 1 fps is generally sufficient for measuring forward movement, though likely captures only part of other major movement behaviors (particularly intermittent reverse
movements—so called “reversals”—whose duration is on the order of 1–2 seconds or less) [22, 33]. For a given minute during the recording session, we compared the total distance traveled by an animal (the sum of all displacements measured from second-by-second imaging) to its net displacement across the first and last images taken over that minute (i.e. the displacement between the first and last image acquired, producing an effective 1 fpm; n = 48 animals × 20 minute intervals = 9600 data points). The overall correlation between net and total displacement is strong with an \( R^2 = 0.60 \) (95% CI 0.50–0.68 generated by bootstrap resampling measurements of random subsets of 43 animals/90% of animals).

Minute-by-minute recordings of early-adult movement reveal early-life movement decline

Because minute-by-minute recordings capture an appreciable fraction of the variability in total second-by-second movement, we used 1 fpm recordings to determine how movement changes during the first days of early adulthood in *C. elegans*. The first 4–6 days of adulthood are often thought of as a period of vigorous activity prior to an overt, qualitative decline in movement rates [7, 19, 21, 25, 34]. Consequently, we chose to focus on this period when assessing patterns of early-adult decline.

We collected movement data from 33 individuals at 1 fpm throughout the first 6 days of adulthood. We observed both substantial variability in movement among individuals, and within single individuals over time. Some individuals showed relatively high and/or sustained periods of maximal movement (Fig 3A, left panel) while others exhibited a relatively lower baseline level of activity (Fig 3A, right panel).
Fig 3. Animal movement shows substantial decline during early-adult life. (A) Examples of movement taken at 1 fpm during the first day of life. Shown are data from two individuals with relatively high (left panel) and low (right panel) movement. (B) Corresponding movement declines across the observation period for the respective individuals in (A), integrated over 3-hour intervals to yield 8 timepoints a day. Rates of decline and coefficients of determination are estimated by linear regression. (C) Change in movement behavior averaged across all individuals during the
We next examined how each individual’s level of activity changed throughout early adulthood. We quantified total movement at 3 hour intervals by summing the total minute-to-minute displacement over each 3 hour interval (Fig 3B) [21, 35], and measured the overall trend in movement during young adulthood with linear regression. The average trend over time, across the whole population, shows an approximately two-fold decline in movement, from ~60 mm per 3-hour block at day 0 of adulthood to less than 30 mm at day 6 (Fig 3C). Moreover, all individuals showed decreasing movement over the first six days of adulthood (Fig 3D and 3E). For most individuals (26/33; ~79%), these declines were statistically distinguishable from zero using the Wald test for linear regression coefficients (Fig 3D). An alternate analysis using bootstrap confidence intervals on the slope of decline showed that none of the 95% confidence intervals included zero (Fig 3D). Overall, these findings suggest that at least a majority of individuals experience substantial declines in movement during the first days of adult life.

Early adult movement is variable among individuals and is not a persistent trait

The observation that movement behavior can decline during early adulthood raises a related question—do all individuals experience a similar pattern of decline? More specifically, how different are individuals’ movement behavior across the first few days of adulthood? On cursory inspection, animals show substantial differences in daily activity even on the first day of adulthood (Fig 3A & 3B). When movement was quantified by total minute-to-minute distance traveled during day 1 of adulthood, total movement spanned a 4-fold range across all individuals (Fig 4A). Expressed as the coefficient of variation (CV; the ratio of the standard deviation to mean of total day movement), the level of variability amounts to ~1/4 (27%) of the average distance that an animal traveled that day. Interestingly, this variability was evident across each day and increased during days 5 and 6 as movement continued to decline (Fig 4A).

Having observed such variability across animals in the setting of significant age-related declines, we questioned whether an animal’s level of movement reflected a persistent trait; that is, do some animals always move more than others, even amid an overall decline in movement with age? We thus asked if movement on any given day was predictive of movement on a future day. Surprisingly, we found that movement among the first 2–3 days of adulthood was not predictive of movement on any other day out to the end of early-adulthood on day 6 (Fig 4B, top panel; Fig 4C). Greater correlations among daily movement scores began to emerge at days 4–5 of adulthood (Fig 4B, bottom panel; Fig 4C). Taken together, these findings suggest that youthful movement is highly variable both between and within individual animals, despite being overlaid on an overall downward trend through early adulthood.

Standard movement assays are underpowered to resolve early adult movement decline

It is curious that, despite extensive study of *C. elegans* movement through aging, early adulthood movement declines are rarely studied and/or reported (with several notable exceptions [19–21]). One potential reason for this is the high degree of between- and within-individual variability in movement noted above. However, we also wondered whether the design of
specific data-acquisition protocols could exacerbate this issue. To answer this question, we performed simulations to study the effects of different measurement protocols on the types of conclusions that can subsequently be drawn.

In these simulation studies, we resampled image data from the collected “gold-standard” data collected at 1 fpm and to simulate the movement scores that would be generated for each day.
individual under several different data-collection protocols. For consistency with the analyses of minute-to-minute movement in previous sections, we summed movement across 3-hour blocks when comparing protocols. We then compared the resulting series of reconstructed movement measurements to those calculated from the original data.

We examined two different measurement protocols, both common in the *C. elegans* literature (Fig 5A). First, we defined a “Single Snapshot” protocol that measures movement as the displacement of an animal between timepoints at a prescribed interval (i.e. displacement in animal position between two images taken at intervals of 2 minutes to almost 3 hours apart) [21]. Second, we defined a "Daily Video" protocol that measures 15 minutes of movement at 1 fps, once a day per individual. This is analogous to standard plate assays in which animals are recorded at high frame rate for a fixed amount of time each day [17, 26, 34, 36]. Unlike the “Single Snapshot” protocol which we can readily examine by subsampling our continuous 1 fpm imaging data, we cannot directly produce a dataset equivalent to the “Daily Video” protocol from the 1 fpm image data we obtained. However, as noted above, over 15-minute intervals, summed displacements from images taken at 1 fps and 1 fpm are very strongly correlated ($R^2 = 0.81$; S1 Fig). Thus, we can effectively simulate “Daily Video” measurements for a given day by summing the minute-to-minute displacement over 15 minutes at the start of that 24 hour period.

We found that the Single Snapshot protocol was unable to recapitulate decline in animals seen with 1 fpm imaging. With increasing duration between subsequent images, an animal’s displacement between images more poorly resembles total 1 fpm movement over time (Fig 5B). Across all timepoints during early life for a given animal, Single Snapshot movement correlates increasingly worse with overall movement as the inter-image interval is increased (Fig 5C). Across all animals, this correlation falls to a moderate correlation at 15-minute inter-image intervals (mean $R^2 = 0.44$), and virtually no correlation at 150-minute intervals (mean $R^2 = 0.03$; Fig 5D). At this last inter-image interval, the Single Snapshot protocol does not allow identification of statistically significant age-related movement decline in any individual (Fig 6). Taken together, these findings suggest that inter-image intervals of 2–5 minutes are most representative of an animal’s overall movement across the lifespan.

The Daily Video protocol had similar difficulty in recapitulating overall movement across this period. Qualitatively, an animal’s total summed movement across a whole day is unlikely to be well-approximated by a single short burst of observations (Fig 5E), unless movement across all fifteen-minute intervals throughout the day are very highly correlated. Reports of periodic/diurnal patterns in *C. elegans* movement behavior [37–39] make this a priori unlikely. And indeed, movement during the specified 15-minute interval correlates at best modestly with each day’s total movement (mean $R^2$ across animals = 0.22; Fig 5F and 5G). Moreover, when used to assess early life movement decline, the Daily Video protocol also fails to identify statistically significant age-related movement decline in any individual (Fig 6). Of note, assessing movement via 15-minute samples twice per day (as done in some studies) yielded a improved correlation between 15-minute movement and (half-day) movement activity (mean $R^2$ across animals = 0.46); however this does not substantially improve the ability to discern early-adult movement declines (S2 Fig).

**Optimization of measurement protocols produces a simplified assay with improved sensitivity for early-adult movement decline**

These results suggest that maximizing the quantity of measurements (number of timepoints) or quality of measurements (data acquisition rate at each timepoint) individually fails to reproduce trends in early-life movement seen with continuous 1 fpm image acquisition. This caused
Fig 5. Comparison of validity of three movement protocol schemes. (A) Graphical cartoon of the various protocols of interest. The “Continuous Sampling” condition is the baseline-quality data collected in this study. In the “Single Snapshot” protocol, one image per timepoint is taken, and the movement score for that timepoint is the displacement of the individual’s centroid from the previous timepoint. The “Daily Video” protocol consists of 15 images obtained per timepoint at 1 fpm; the movement score for that timepoint is the summed total displacement across all 15 images acquired in that day’s timepoint. Finally, in the “Hybrid Multipass” protocol, several images per timepoint are taken with a wider temporal spacing (allowing image acquisition for several individuals to be interleaved) and the total displacement across those images is used as the movement score for that timepoint. Here, we acquire 7 images each spaced 3 minutes apart (for a total interval of 18 minutes). (B-D) Illustration of data generated by the Single Snapshot protocol at various inter-timepoint intervals. (B) One individual’s activity across the first 6 days of adulthood. (C) For that individual, the correlation between the total displacement during those intervals (measured from gold-standard 1 fpm data) and net displacement using the specified inter-timepoint interval. (D) Kernel density plot showing the distribution of correlations between 1 fps and Single Snapshot obtained for all individuals at the various inter-timepoint intervals. (E-G) Illustration of the data generated by the Daily Video protocol at various inter-timepoint intervals. (E) Activity from the same individual in panel B, across the first 6 days of adulthood. (F) The correlation between the total displacement during
us to wonder whether a protocol with measurements of intermediate frequency and quality might fare better.

We thus evaluated a “Hybrid Measurement” protocol that combines sparser imaging over a relatively long duration of acquisition. This protocol was motivated by the observation that movement on the order of a few (1–6 minutes) minutes is well-correlated to second-scale movement (e.g. Fig 5B). As a proof of principle, we used this observation to maximize the quality of measurements for our group’s Worm Corral system. Specifically in our workflow, separately housed individuals on up to 6–7 devices are serially imaged on an upright compound microscope at high-magnification with a typical throughput of ~600–700 animals with a ~3 hour inter-timepoint interval. On average, moving to a new individual on a device, automatically focusing, and acquiring an image takes ~1.4 sec. Under these constraints, we devised a protocol in which movement is measured as total displacement across seven images acquired over 18 minutes total (yielding an inter-image interval of three minutes). This protocol preserves the typical throughput of our system while allowing multiple timepoint acquisitions per day of movement across several minutes.

When we simulated this protocol, it displayed marked improvement in measuring movement and age-related decline. Individual trajectories of decline were more qualitatively similar to those from minute-to-minute movement measurements compared to those estimated by the other protocols (cf. Fig 5B and 5H). Additionally, movement measurements using this protocol were correlated with total movement measured from 1 fpm imaging over the total 3-hour inter-timepoint interval (mean $R^2 = 0.44$; Fig 5I and 5J). Most importantly, we found

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<th>Protocol</th>
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<td>Continuous Sampling</td>
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<td>Daily Video Protocol</td>
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<td>Hybrid Protocol</td>
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Fig 6. Visual comparison of measurable decline in activity across different image acquisition protocols. Activity traces are labeled for all 33 animals and colored red vs. green to denote non-significant vs. significant decline, respectively (ordering of animals as in Fig 3E).

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that measurement of movement decline using this scheme was able to identify statistically significant declines in just over half of animals that had showed significant declines at a resolution of 1 fpm (14/26). These findings suggest that a simple modification to movement measurement that allows for frequent sampling of moderate-quality measurements may be sufficiently sensitive to capture early-life movement decline.

**Discussion**

In this study, we questioned whether young adult individuals are subject to age-related declines in movement analogous to those seen in older individuals. We subsequently found evidence that movement activity begins to decline as early as the first days of adulthood. Moreover, early-adult movement behavior is highly variable across individuals; in fact, the level of variation in total movement on this first day of adulthood is comparable to or higher than the variation typically seen in other life traits in wild-type *C. elegans*, including lifespan [21] and fecundity [40]. Taken together, these findings suggest that the vigorous movement that characterizes early adult life is not some generic state maintained consistently over time and across all individuals until some qualitative failure point. Rather, movement is a relative trait that appears to generally decline over time in early adulthood, even in otherwise “healthy” animals, and at a rate that is nevertheless quite variable across individuals.

It is important to note that our data do not conclusively demonstrate whether or not there are individuals that truly show no appreciable decline in movement over this span. The subset of individuals we studied that did not have statistically significant declines in movement (based on a parametric test of the regression coefficients) still all had negative point estimates of the slope in movement over time (indicating modest declines in movement rate), with bootstrapped 95% confidence intervals around those point estimates that did not include zero. Such a discrepancy reflects no strong statistical evidence as to the extent of decline in this subset of animals.

That individuals show such variability in early-life movement also suggests that there may exist interesting biology of individual differences in movement behavior, which remains as-yet largely unexplored. In many studies, average movement has a nearly constant and relatively constrained range of values representing “healthy” young-adult movement that only declines starting at days 6–8. We suspect that this high level of activity reflects a ceiling effect of the movement measurements used. That is, our results suggest that measurements based on brief intervals of observed activity may not have the sensitivity to detect inter- and intraindividual differences in movement during the first several days of adulthood. Indeed, a recent study of interindividual variability in movement during development using high-temporal resolution (3 fps) also suggests significant variation in movement activity at least on the first day of adulthood [41].

The experimental design for this study made use of the observation that the distance traveled when displacements are measured second-to-second is well-approximated by the distance traveled when displacements are captured sixty times less frequently, with minute-to-minute measurements (\(R^2 \sim 0.5–0.68\) on a minute-to-minute basis and upwards of 0.81 at longer timescales). This is supported by corroborating evidence from other studies suggesting that *C. elegans* movement is correlated across multiple timescales [42, 43]. Consequently, we believe that measuring movement at minute timescales is representative of bulk animal movement, though we caution that it may well not be representative of fine animal movement—e.g. turning, reversals, or head turning.

Motivated by these findings, we identified an intermediate measurement regime that provides a substantial improvement in sensitivity to early-adult movement. In particular, we find that the advantage of sampling movement more often over the day more than outweighs the
disadvantages of conducting that sampling at a coarser time-scale (on the order of minutes). We hope that these results will inspire others to determine whether optimized movement measurement protocols in different culture systems can also be as or more accurate than current protocols while also requiring fewer images to be analyzed overall.

It is important to note several unique features of our culture system that may limit the direct generalizability of these results to all settings. First, the “worm corral” system involves a relatively confined environment (a roughly circular bacterial food pad ≈1.5–1.6 mm, or ~1.5–1.75 typical body lengths, in diameter). This smaller area might influence behavior, as it is known animals on food tend to be more quiescent than those off food [39], potentially leading to underestimation of an animal’s overall capacity to move [17, 39]. Moreover, the small food pad size places a ceiling over the maximum displacement measurable between timepoints, as total movement distance on the scale of minutes or more is often larger than the size of the pad. This is consistent with our observations that inter-imaging intervals greater than 5–10 minutes compromise movement measurements. In culture systems with larger movement arenas, longer inter-imaging intervals may be as or more useful than those employed here.

Another feature of our culture system is the use of relatively high magnification compared to other long-term imaging apparatuses, which requires moving the stage to and focusing on individual animals one after another. In this context, our hybrid protocol provides the advantage that imaging bouts for one individual can be interleaved with imaging for others, increasing overall throughput. Optimal protocols may differ for other longitudinal observation systems in which many identifiable individuals can be simultaneously observed at once [25, 44].

Last, we examined unstimulated movement only, which may be less sensitive for observing age-related changes across life [25, 34]. Nevertheless, our finding of modest early-life declines in movement ability and/or propensity in most individuals provides a similar opportunity for researchers with different systems to calibrate and optimize their own motion-measurement protocols. Likewise, we suspect that the basic scheme of the optimized “hybrid” protocol we identified will be of use to others: the combination of moderately sensitive measurements repeated multiple times daily may be both more efficacious than infrequent, high-sensitivity imaging bouts and also allow greater throughput than continuous high-sensitivity imaging.

Supporting information
S1 Fig. Correlation between second-scale and minute-scale total movement over 15 minute intervals. Comparison of total displacement at 1 fps over 15 minutes and the summed net displacement at 1 fpm over the same 15-minute interval (n = 48 animals) using the pilot 1 fps dataset. The overall correlation between total and net summed displacement is $R^2 = 0.81$ (95% CI 0.67–0.90 generated by bootstrap resampling the measurements of random subsets of 43 animals/90% of animals).
(SPS)

S2 Fig. Performance of twice-daily video assessment of movement. (A) The same individual in Fig 5A and 5E’s activity across all 6 days of life using a simulated twice-daily video measurement. (B) The correlation between the total displacement during the sampling period and the subsequent time (i.e. given by the intertimepoint interval). (C) Kernel density plot showing the distribution of correlations obtained for all individuals with the specified intertimepoint intervals.
(SPS)
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