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DeepFly3D: A deep learning-based approach for 3D limb and appendage tracking in tethered, adult *Drosophila*

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¹⁰ **Abstract** Studying how neural circuits orchestrate limbed behaviors requires the precise

- measurement of the positions of each appendage in 3-dimensional (3D) space. Deep neural
- networks can estimate 2-dimensional (2D) pose in freely behaving and tethered animals. However,
- the unique challenges associated with transforming these 2D measurements into reliable and
- ¹⁴ precise 3D poses have not been addressed for small animals including the fly, *Drosophila*
- *melanogaster*. Here we present DeepFly3D, a software that infers the 3D pose of tethered, adult
- ¹⁶ *Drosophila*—or other animals—using multiple camera images. DeepFly3D does not require manual
- calibration, uses pictorial structures to automatically detect and correct pose estimation errors, and
- uses active learning to iteratively improve performance. We demonstrate more accurate
- ¹⁹ unsupervised behavioral embedding using 3D joint angles rather than commonly used 2D pose
- data. Thus, DeepFly3D enables the automated acquisition of behavioral measurements at an
- ²¹ unprecedented level of resolution for a variety of biological applications.
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23 Introduction

²⁴ The precise quantification of movements is critical for understanding how neurons, biomechanics,

- ²⁵ and the environment influence and give rise to animal behaviors. For organisms with skeletons and
- ²⁶ exoskeletons, these measurements are naturally made with reference to 3D joint and appendage
- ²⁷ locations. Paired with modern approaches to simultaneously record the activity of neural popu-
- lations in tethered, behaving animals (Dombeck et al., 2007; Seelig et al., 2010; Chen et al., 2018),
- ²⁹ 3D joint and appendage tracking promises to accelerate the dissection of neural control principles,
- particularly in the genetically tractable and numerically simple nervous system of the fly, *Drosophila melanogaster*.
- However, algorithms for reliably estimating 3D pose in such small animals have not yet been developed. Instead, multiple alternative approaches have been taken. For example, one can affix and use small markers—reflective, colored, or fluorescent particles—to identify and reconstruct
- keypoints from video data (*Bender et al., 2010; Kain et al., 2013; Todd et al., 2017*). Although this
- ³⁶ approach works well on humans (*Moeslund and Granum, 2000*), in smaller animals markers likely
- hamper movements, are difficult to mount on sub-millimeter scale limbs, and, most importantly,
- measurements of one or even two markers on each leg (*Todd et al., 2017*) cannot fully describe
- ³⁹ 3D limb kinematics. Another strategy has been to use computer vision techniques that operate
- 40 without markers. However, these measurements have been restricted to 2D pose in freely behaving

animals. Before the advent of deep learning, this was accomplished by matching the contours of 41 animals seen against uniform backgrounds (Mori and Malik, 2006), measuring limb tip positions 42 using complex TIRF-based imaging (Mendes et al., 2013), or measuring limb segments using active 43 contours (Uhlmann et al., 2017). In addition to being limited to 2D rather than 3D pose, these meth-44 ods are complex, time-consuming, and error-prone in the face of long data sequences, cluttered 45 backgrounds, fast motion, and occlusions that naturally occur when animals are observed from a 46 single 2D perspective. 47 As a result, in recent years the computer vision community has largely forsaken these techniques 48 in favor of deep learning-based methods. Consequently, the effectiveness of monocular 3D human 49 pose estimation algorithms has improved greatly. This is especially true when capturing human 50 movements for which there is enough annotated data to train deep networks effectively. Walking 51 and upright poses are prime examples of this, and state-of-the-art algorithms (Paylakos et al., 52 2017a: Tome et al., 2017: Popa et al., 2017: Moreno-noguer, 2017: Martinez et al., 2017: Mehta 53 et al., 2017; Rogez et al., 2017; Pavlakos et al., 2017b; Zhou et al., 2017; Tekin et al., 2017; Sun 54 et al., 2017) now deliver impressive real-time results in uncontrolled environments. Increased 55 robustness to occlusions can be obtained by using multi-camera setups (Elhayek et al., 2015; 56 Rhodin et al., 2016: Simon et al., 2017: Paylokos et al., 2017b) and triangulating the 2D detections. 57 This improves accuracy while making it possible to eliminate false detections. 58 These advances in 2D pose estimation have also recently been used to measure behavior in 59 laboratory animals. For example, DeepLabCut provides a user-friendly interface to DeepCut, a 60 state-of-the-art human pose estimation network (Mathis et al., 2018), and LEAP (Pereira et al., 61 2019) can successfully track limb and appendage landmarks using a shallower network. Still, 2D 62 pose provides an incomplete representation of animal behavior: important information can be lost 63 due to occlusions, and movement quantification is heavily influenced by perspective. Unfortunately, 64 techniques used to translate human 2D pose to 3D pose cannot be easily transferred for the study 65 of small animals like Drosophila: adult flies are approximately 2.5 mm long, have many appendages 66 and joints, are translucent, and in most laboratory experiments are only illuminated using infrared 67 light (to avoid visual stimulation)—precluding the exploitation of color information. Moreover, 68 precisely registering multiple camera viewpoints using traditional approaches would require the fabrication of a prohibitively small checkerboard pattern, along with the tedious labor of repeatedly 70 calibrating using a small, external target. 71 To overcome these challenges, we introduce DeepFlv3D, a deep learning-based software pipeline 72 that achieves comprehensive, rapid, and reliable 3D pose estimation in tethered, behaving adult 73 Drosophila (Figure 1, Figure 1-video 1). DeepFlv3D is applied to synchronized videos acquired 74 from multiple cameras (Figure 12). It first uses a state-of-the-art deep network (Newell et al., 75 2016) and then enforces consistency across views (Figure 8). This makes it possible to eliminate 76 spurious detections, achieve high 3D accuracy, and use 3D pose errors to further fine-tune the 77 deep network to achieve even better accuracy (*Figure 2*). To register the cameras. DeepFlv3D uses 78 a novel calibration mechanism in which the fly itself is the calibration target (*Figure 7*). Thus, the 79 user doesn't need to manufacture a prohibitively small calibration pattern, or repeat cumbersome 80 calibration protocols. We explain how users can modify the codebase to extend DeepFly3D for 81 3D pose estimation in other animals (Figure 11 and see Methods). Finally, we demonstrate that 82 unsupervised behavioral embedding of 3D joint angle data (Figure 4) is robust against problematic 83 artifacts present in embeddings of 2D pose data (*Figure 3*). In short, DeepFlv3D delivers 3D pose 84

estimates reliably, accurately, and with minimal manual intervention while also providing a critical

⁸⁶ tool for automated behavioral data analysis.

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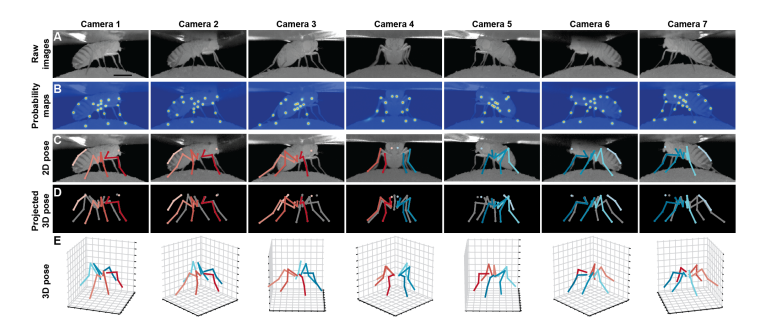


Figure 1. Deriving 3D pose from multiple camera views. **(A)** Raw image inputs to the Stacked Hourglass deep network. **(B)** Probability maps output from the trained deep network. For visualization purposes, multiple probability maps have been overlaid for each camera view. **(C)** 2D pose estimates from the Stacked Hourglass deep network after applying pictorial structures and multi-view algorithms. **(D)** 3D pose derived from combining multiple camera views. For visualization purposes, the 3D pose has been projected onto the original 2D camera perspectives. **(E)** 3D pose rendered in 3D coordinates. Immobile thorax-coxa joints and antennal joints have been removed for clarity. The following video supplement is available for this figure:

Figure 1-video 1. Deriving 3D pose from multiple camera views during backward walking in an optogenetically stimulated MDN>CsChrimson fly. https://drive.google.com/file/d/15nGQRgrjY4dyGh0GFr5eZrRQuOR6Z4fK/view?usp=sharing.

87 **Results**

88 DeepFly3D

The input to DeepFly3D is video data from seven cameras (*Figure 12*). These images are used to identify the 3D positions of 38 landmarks per animal: (i) five on each limb – the thorax-coxa, coxa-femur, femur-tibia, and tibia-tarsus joints as well as the pretarsus, (ii) six on the abdomen - three on each side, and (iii) one on each antenna - useful for measuring head rotations. Our software incorporates a number of innovations designed to ensure automated, high-fidelity, and reliable 3D pose estimation:

 Calibration without external targets: Estimating 3D pose from multiple images requires 95 calibrating the cameras to achieve a level of accuracy that is commensurate with the tar-96 get size—a difficult challenge when measuring leg movements for an animal as small as 97 Drosophila. Therefore, instead of using a typical external calibration grid, DeepFly3D uses 98 the fly itself as a calibration target. It detects arbitrary points on the fly's body and relies on 99 bundle-adjustment (Chavdarova et al., 2018) to simultaneously assign 3D locations to these 100 points and to estimate the positions and orientations of each camera (Figure 7). To increase 101 robustness, it enforces geometric constraints that apply to tethered flies with respect to limb 102 segment lengths and ranges of motion. 103 **Geometrically consistent reconstructions:** Starting with a state-of-the-art deep network 104

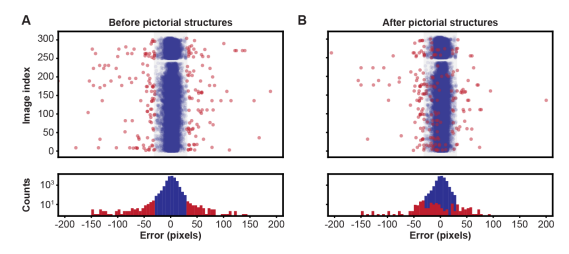
for 2D keypoint detection in individual images (*Newell et al., 2016*), DeepFly3D enforces geometric consistency constraints across multiple synchronized camera views. When triangulating 2D detections to produce 3D joint locations, it relies on pictorial structures and belief propagation message passing (*Felzenszwalb and Huttenlocher, 2005*) to detect and further correct erroneous pose estimates (*Figure 8*).

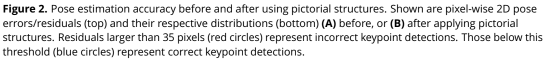
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- Self-supervision and active learning: We also use multiple view geometry as a basis for active learning. Thanks to the redundancy inherent in obtaining multiple views of the same animal, we can detect erroneous 2D predictions for correction (*Figure 10*) that would most efficiently train the 2D pose deep network. This approach greatly reduces the need for time-consuming manual labeling (*Simon et al., 2017*). We also use pictorial structure corrections to
- fine-tune the 2D pose deep network. Self-supervision constitutes 85% of our training data.

Improving 2D pose using pictorial structures and active learning

We validated our approach using a challenging dataset of 2063 frames manually annotated using 117 the DeepFlv3D annotation tool (*Figure 6*) and sampled uniformly from each camera. Images for 118 testing and training are 480×960 pixels. The test dataset included challenging frames and occasional 119 motion blur to increase the difficulty of pose estimation. For training, we used a final training 120 dataset of 37,000 frames, an overwhelming majority of which were first automatically corrected 121 using pictorial structures. On test data, we achieved a Root Mean Square Error (RMSE) of 13.9 122 pixels. Setting a 50 pixel threshold for PCK (percentage of correct keypoints) computation, we 123 observed a 98.2% general accuracy before applying pictorial structures. We found that application 124 of pictorial structures corrected 59% of erroneous predictions, increasing the final accuracy to 125 99.2%. These improvements are illustrated in *Figure 2*. Pictorial structure failures were often due 126 to pose ambiguities resulting from heavy motion blur. These remaining errors were automatically 127 detected with multi-view redundacy using *Equation 6*, and earmarked for manual correction using 128 the DeepFly3D GUI (Figure 9). 129





3D pose permits robust unsupervised behavioral classification

Unsupervised behavioral classification approaches enable the unbiased quantification of animal 131 behavior by processing data features—image pixel intensities (Berman et al., 2014: Cande et al., 132 2018), limb markers (Todd et al., 2017), or 2D pose (Pereira et al., 2019)—to cluster similar behav-133 ioral epochs without user intervention and to automatically distinguish between otherwise similar 134 actions. However, with this sensitivity may come a susceptibility to features unrelated to behavior. 135 These may include changes in image size or perspective resulting from differences in camera angle 136 across experimental systems, variable mounting of tethered animals, and inter-animal morpho-137 logical variability. In theory, each of these issues can be overcome—providing scale and rotational 138 invariance—by using 3D joint angles rather than 2D pose for unsupervised embedding. 139

To test this possibility, we performed unsupervised behavioral classification on video data 140 taken during optogenetic stimulation experiments that repeatedly and reliably drove specific ac-141 tions. Specifically, we optically activated CsChrimson (Chen et al., 2013) to elicit backward walking 142 in MDN>CsChrimson animals (Figure 4-video 1) (Bidave et al., 2014), or antennal grooming in 143 aDN>CsChrimson animals (Figure 4-video 2) (Hampel et al., 2015). We also stimulated control 144 animals lacking the UAS-CsChrimson transgene (Figure 4-video 3)(MDN-GAL4/+ and aDN-GAL4/+). 145 First, we performed unsupervised behavioral classification using 2D pose data from three adia-146 cent cameras containing keypoints for three limbs on one side of the body. Using these data, we 147 generated a behavioral map (Figure 3A). In this map each individual cluster would ideally repre-148 sent a single behavior (e.g., backward walking, or grooming) and be populated by nearly equal 149 amounts of data from each of the three cameras. This was not the case: data from each camera 150 covered non-overlapping regions and clusters (*Figure 3B-D*). This effect was most pronounced 151 when comparing regions populated by cameras 1 and 2 versus camera 3. Therefore, because the 152 underlying behaviors were otherwise identical (data across cameras were from the same animals 153 and experimental time points), we concluded that unsupervised behavioral classification of 2D pose 154 data is highly sensitive to corruption by viewing angle differences. 155

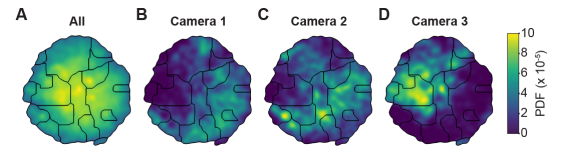


Figure 3. Unsupervised behavioral classification of 2D pose data is sensitive to viewing angle. **(A)** Behavioral map derived using 2D pose data from three adjacent cameras (Cameras 1, 2, and 3) but the same animals and experimental time points. Shown are clusters (black outlines) with enriched (yellow), or sparsely (blue) populated data. Different clusters are enriched for data from either **(B)** camera 1, **(C)** camera 2, or **(D)** camera 3. Behavioral embeddings were derived using 1 million frames during 4 s of optogenetic stimulation of MDN>CsChrimson (n=6 flies, n=29 trials), aDN>CsChrimson (n=6 flies, n=30 trials), and wild-type control animals (MDN-GAL4/+: n=4 flies, n=20 trials. aDN-GAL4/+: n=4 flies, n=23 trials).

By contrast, performing unsupervised behavioral classification using DeepFly3D-derived 3D joint angles resulted in a map (*Figure 4*) with a clear segregation and enrichment of clusters for different GAL4 drivers lines and their associated behaviors (i.e., backward walking (*Figure 4-video 4*), grooming (*Figure 4-video 5*), and forward walking (*Figure 4-video 6*)). Thus, 3D pose overcomes serious issues arising from unsupervised embedding of 2D pose data, enabling more reliable and robust behavioral data analysis.

162 **Discussion**

We have developed DeepFly3D, a deep learning-based 3D pose estimation system that is optimized 163 for quantifying limb and appendage movements in tethered, behaving *Drosophila*. By using multiple 164 synchronized cameras and exploiting multi-view redundancy, our software delivers robust and accu-165 rate pose estimation at the sub-millimeter scale. Our approach relies on supervised deep learning 166 to train a neural network that detects 2D joint locations in individual camera images. Importantly, 167 our network becomes increasingly competent as it runs: By leveraging the redundancy inherent 168 to a multiple-camera setup, we iteratively reproject 3D pose to automatically detect and correct 169 2D errors, and then use these corrections to further train the network without user intervention. 170 Ultimately, we may work solely with monocular images by lifting the 2D detections (Pavlakos et al., 171 2017b) to 3D or by directly regressing to 3D (Tekin et al., 2017) as has been achieved in human pose 172

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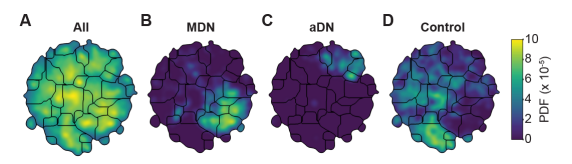


Figure 4. Unsupervised behavioral classification of 3D joint angle data. Behavioral embeddings were calculated using 3D joint angles from the same 1 million frames used in Figure 3. **(A)** Behavioral map combining all data during 4 s of optogenetic stimulation of MDN>CsChrimson (n=6 flies, n=29 trials), aDN>CsChrimson (n=6 flies, n=30 trials), and wild-type control animals (For MDN-Gal4/+, n=4 flies, n=20 trials. For aDN-Gal4/+ n=4 flies, n=23 trials). The same behavioral map is shown with only the data from **(B)** MDN>CsChrimson stimulation, **(C)** aDN>CsChrimson stimulation, or **(D)** control animal stimulation. Associated videos reveal that these distinct map regions are enriched for backward walking, antennal grooming, and forward walking, respectively. The following video supplements are available:

Figure 4-video 1. Representative MDN CsChrimson optogenetically activated backward walking. https://drive.google.com/file/d/1YY98bo2ZbjLotyiTHdViey5zfhKow4Jx/view?usp=sharing Figure 4-video 2. Representative aDN>CsChrimson optogenetically activated antennal grooming. https://drive.google.com/file/d/1_QBgt7P6DhR9hHkNArQIOyNaZALTQumk/view?usp=sharing Figure 4-video 3. Representative control animal behavior during illumination. https://drive.google.com/file/d/1OolwMCSyZFyJ6TQ6sTlcJIaMCT69JKH2/view?usp=sharing Figure 4-video 4. Sample behaviors from 3D pose cluster enriched in backward walking. https://drive.google.com/file/d/1H-R1PmcusV55Yw7c_4dKVFaGtJM-FG9M/view?usp=sharing Figure 4-video 5. Sample behaviors from 3D pose cluster enriched in antennal grooming. https://drive.google.com/file/d/1f7TaF8FTWNwuvpdK9hV0IX7tt6f2QjXo/view?usp=sharing Figure 4-video 6. Sample behaviors from 3D pose cluster enriched in forward walking. https://drive.google.com/file/d/1f7TaF8FTWNwuvpdK9hV0IX7tt6f2QjXo/view?usp=sharing Figure 4-video 6. Sample behaviors from 3D pose cluster enriched in forward walking.

estimation studies. In the Methods section, we explain in detail how organism-specific features of DeepFly3D—bone segment length, number of legs, and camera focal distance—can be modified to

175 study, for example, humans (*Figure 11*), primates, rodents, or other insects.

As in the past, we anticipate that the development of new technologies for quantifying behavior 176 will open new avenues and enhance existing lines of investigation. For example, deriving 3D 177 pose using DeepFly3D can improve the resolution of studies examining how neuronal stimulation 178 influences animal behavior (Cande et al., 2018: McKellar et al., 2019), the precision and predictive 179 power of efforts to define natural action sequences (Seeds et al., 2014: McKellar et al., 2019), the 180 assessment of interventions that target models of human disease (Feany and Bender, 2000; Hewitt 181 and Whitworth, 2017), and the linking of neural activity with animal behavior—when coupled with 182 recording technologies like 2-photon microscopy (Seelig et al., 2010; Chen et al., 2018). Importantly, 183 3D pose dramatically improves the robustness of unsupervised behavioral classification approaches. 184 Therefore, DeepFly3D is a critical step toward the ultimate goal of achieving fully-automated, high-185 fidelity behavioral data analysis. 186

187 Materials and Methods

188 With synchronized Drosophila video sequences from seven cameras in hand, the first task for

- 189 DeepFly3D is to detect the 2D location of 38 landmarks. These 2D locations of the same landmarks
- ¹⁹⁰ seen across multiple views are then triangulated to produce 3D pose estimates. This pipeline is
- depicted in *Figure 5*. First, we will describe our deep learning-based approach to detect landmarks
- ¹⁹² in images. Then, we will explain the triangulation process that yields full 3D trajectories. Finally, we

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¹⁹³ will describe how we identify and correct erroneous 2D detections automatically.

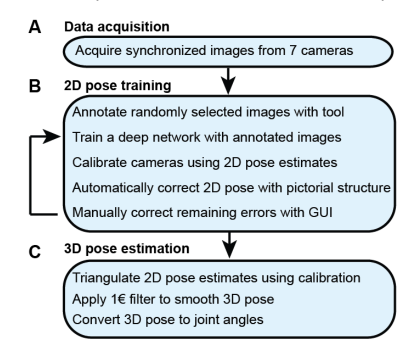


Figure 5. The DeepFly3D pose estimation pipeline. **(A)** Data acquisition from the multi-camera system. **(B)** Training and retraining of 2D pose. **(C)** 3D pose estimation.

Deep Network Architecture. We aim to detect five joints on each limb, six on the abdomen. 194 and one on each antenna, giving a total of 38 keypoints per time instance. To achieve this, we 195 adapted a state-of-the-art Stacked Hourglass human pose estimation network (Newell et al., 2016) 196 by changing the input and output layers to accommodate a new input image resolution and a 197 different number of tracked points. A single hourglass stack consists of residual bottleneck modules 198 with max pooling, followed by up-sampling layers and skip connections. The first hourglass network 199 begins with a convolutional layer and a pooling layer to reduce the input image size from 256×512 200 to 64×128 pixels. The remaining hourglass input and output tensors are 64×128 . We used 8 stacks 201 of hourglasses in our final implementation. The output of the network is a stack of probability 202 maps, also known as heatmaps or confidence maps. Each probability map encodes the location 203 of one keypoint, as the belief of the network that a given pixel contains that particular tracked 204 point. However, probability maps do not formally define a probability distribution: their sum over 205 all pixels does not equal 1. 206

2D pose training dataset. We trained our network for 19 keypoints, resulting in the tracking 207 of 38 points when both sides of the fly are accounted for. Determining which images to use for 208 training purposes is critical. The intuitively simple approach—training with randomly selected 209 images—may lead to only marginal improvements in overall network performance. This is because 210 images for which network predictions can already be correctly made give rise to only small gradients 211 during training. On the other hand, manually identifying images that may lead to incorrect network 212 predictions is highly laborious. Therefore, to identify such challenging images, we exploited the 213 redundancy of having multiple camera views (see section 3D pose correction). Outliers in individual 214 camera images were corrected automatically using images from other cameras, and frames that still 215 exhibited large reprojection errors on multiple camera views were selected for manual annotation 216 and network retraining. This combination of self supervision and active learning permits faster 217 training using a smaller manually annotated dataset (Simon et al., 2017). The full annotation 218 and iterative training pipeline is illustrated in *Figure 5*. In total, 40.063 images were annotated: 219

5,063 were labeled manually in the first iteration, 29,000 by automatic correction, and 6,000 by
 manually correcting those proposed by the active learning strategy.

Deep network training procedure. We trained our Stacked Hourglass network to regress from 222 256 x 512 pixel gravscale video images to multiple 64 x 128 probability maps. Specifically, during 223 training and testing, networks output a $19 \times 64 \times 128$ tensor; one 64×128 probability map per 224 tracked point. During training, we created probability maps by embedding a 2D Gaussian with 225 mean at the ground-truth point and 1px symmetrical extent, i.e., with $\sigma = 1px$ on the diagonal of 226 the covariance matrix. We calculated the loss as the L_2 distance between the ground-truth and 227 predicted probability maps. During testing, the final network prediction for a given point was the 228 probability map pixel with maximum probability. We started with a learning rate of 0.0001 and 229 then multiplied the learning rate by a factor of 0.1 once the loss function plateaued for more than 230 5 epochs. We used an RMSPROP optimizer for gradient descent, following the original Stacked 231 Hourglass implementation, with a batch-size of 8 images. Using 37,000 training images, the Stacked 232 Hourglass network usually converges to a local minimum after 100 epochs (20 hours on a single 233 GPU). 234

Network training details. Variations in each fly's position across experiments are handled 235 by the translational invariance of the convolution operation. In addition, we artificially augment 236 training images to improve network generalization for further image variables. These variables 237 include (i) illumination conditions – we randomly changed the brightness of images using a gamma 238 transformation. (ii) scale – we randomly rescaled images between 0.80x - 1.20x, and (iii) rotation – we 230 randomly rotated images and corresponding probability maps $\pm 15^{\circ}$. This augmentation was enough 240 to compensate for real differences in the size and orientation of tethered flies across experiments. 241 Furthermore, as per general practice, the mean channel intensity was subtracted from each input 242 image to distribute annotations symmetrically around zero. We began network training using 243 pretrained weights from the MPII human pose dataset (Andriluka et al., 2014). This dataset consists 244 of more than 25,000 images with 40,000 annotations, possibly with multiple ground-truth human 245 pose labels per image. Starting with a pretrained network results in faster convergence. However, in 246 our experience, this does not affect final network accuracy in cases with a large amount of training 247 data. We split the dataset into 37,000 training images, 2,063 testing images, and 1,000 validation 248 images. None of these subsets shared common images or common animals, to ensure that 249 the network could generalize across animals, and experimental setups. 5.063 of our training 250 images were manually annotated, and the remaining data were automatically collected using belief 251 propagation, graphical models, and active learning, (see section 3D pose correction). Deep neural 252 network parameters need to be trained on a dataset with manually annotated ground-truth key 253 point positions. To initialize the network, we collected annotations using a custom multicamera 254 annotation tool that we implemented in JavaScript using Google Firebase (*Figure 6*). The DeepFlv3D 255 annotation tool operates on a simple web-server, easing the distribution of annotations across 256 users and making these annotations much easier to inspect and control. We provide a GUI to 257 inspect the raw annotated data and to visualize the network's 2D pose estimation (Figure 9). 258

Computing hardware and software. We trained our model on a desktop computing work station running on an Intel Core i9-7900X CPU, 32 GB of DDR4 RAM, and a GeForce GTX 1080.
 With 37,000 manually and automatically labeled images, training takes nearly 20 hours on a single
 GeForce GTX 1080 GPU. Our code is implemented with Python 3.6, Pytorch 0.4 and CUDA 9.2.

Accuracy analysis. Consistent with the human pose estimation literature, we report accuracy 263 as Percentage of Correct Keypoints (PCK) and Root Mean Squared Error (RMSE). PCK refers to 264 the percentage of detected points lying within a specific radius from the ground-truth label. We 265 set this threshold as 50 pixels, which is roughly one third of the femur-tibia segment. The final 266 estimated position of each keypoint was obtained by selecting the pixel with the largest probability 267 value on the relevant probability map. We compared DeepFlv3D's annotations with manually 268 annotated ground-truth labels to test our model's accuracy. For RMSE, we report the square root 269 of average pixel distance between the prediction and the ground-truth location of the tracked 270

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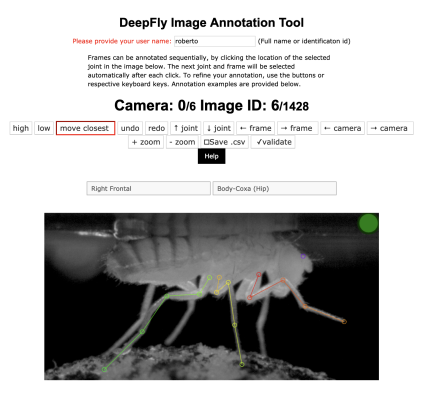


Figure 6. The DeepFly3D annotation tool. This GUI allows the user to manually annotate joint positions on images from each of 7 cameras. Because this tool can be accessed from a web browser, annotations can be performed in a distributed manner across multiple users more easily. A full description of the annotation tool can be found in the online documentation: https://github.com/NeLy-EPFL/DeepFly3D

²⁷¹ point. We remove trivial points such as the body-coxa and coxa-femur—which remain relatively

²⁷² stationary—to fairly evaluate our algorithms and to prevent these points from dominating our

273 accuracy measurements.

274 From 2D landmarks to 3D trajectories

In the previous section, we described our approach to detect 38 2D landmarks. Let $\mathbf{x}_{c,j} \in \mathbb{R}^2$ denote the 2D position of landmark *j* and the image acquired by camera *c*. For each landmark, our task is now to estimate the corresponding 3D position, $\mathbf{X}_j \in \mathbb{R}^3$. To accomplish this, we used triangulation and bundle-adjustment (*Hartley and Zisserman, 2000*) to compute 3D locations, and we used pictorial structures (*Felzenszwalb and Huttenlocher, 2005*) to enforce geometric consistency and to eliminate potential errors caused by misdetections. We present these steps below.

Pinhole camera model. The first step is to model the projection operation that relates a specific \mathbf{X}_j to its seven projections in each camera view $\mathbf{x}_{c,j}$. To make this easier, we follow standard practice and convert all Cartesian coordinates $[x_c, y_c, z_c]$ to homogeneous ones $[x_h, y_h, z_h, s]$ such that $x_c = x_h/s$, $y_c = y_h/s$, $z_c = z_h/s$. From now on, we will assume that all points are expressed in homogeneous coordinates and omit the *h* subscript. Assuming that these coordinates are expressed in a coordinate system whose origin is in the optical center of the camera and whose z-axis is its optical axis, the 2D image projection [u, v] of a 3D homogeneous point [x, y, z, 1] can be bioRxiv preprint first posted online May. 20, 2019; doi: http://dx.doi.org/10.1101/640375. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a CC-Bpeep http://gb02019 attonal license.

288 written as

$$u = U/W,$$

$$v = V/W,$$

$$\begin{bmatrix} U\\ V\\ W \end{bmatrix} = \mathbf{K} \begin{bmatrix} x\\ y\\ z\\ 1 \end{bmatrix}, \text{ with } \mathbf{K} = \begin{bmatrix} f_x & 0 & c_x & 0\\ 0 & f_y & c_y & 0\\ 0 & 0 & 1 & 0 \end{bmatrix},$$
(1)

where the 3 × 4 matrix **K** is known as the *intrinsic parameters matrix*—scaling in the *x* and *y* direction and image coordinates of the principal point c_x and c_y —that characterizes the camera settings.

In practice, the 3D points are not expressed in a coordinate system tied to the camera, especially in our application where we use seven different cameras. Therefore, we use a world coordinate system that is common to all cameras. For each camera, we must therefore convert 3D coordinates expressed in this world coordinate system to camera coordinates. This requires rotating and translating the coordinates to account for the position of the camera's optical center and its orientation. When using homogeneous coordinates, this is accomplished by multiplying the coordinate vector by a 4 × 4 *extrinsic parameters matrix*

$$\mathbf{M} = \begin{bmatrix} \mathbf{R} & \mathbf{T} \\ 0 & 1 \end{bmatrix},\tag{2}$$

where **R** is a 3×3 rotation matrix and **T** a 3×1 translation vector. Combining *Equation 1* and *Equation 2* yields

$$u = U/W,$$

$$v = V/W,$$

$$\begin{bmatrix} U \\ V \\ W \end{bmatrix} = \mathbf{P} \begin{bmatrix} x \\ y \\ z \\ 1 \end{bmatrix}, \text{ where } \mathbf{P} = \mathbf{M}\mathbf{K} \text{ is a } 3 \times 4 \text{ matrix.}$$
(3)

Camera distortion. The pinhole camera model described above is an idealized one. The projections of real cameras deviate from it and these deviations are referred to as distortions and need to be accounted for. The most significant one is known as radial distortion because the error grows with the distance to the image center. For the cameras we use, radial distortion can be expressed as

$$u_{\text{pinhole}} = u \left(1 + k_1^x r^2 + k_2^x r^4 \right) , \qquad (4)$$

$$v_{\text{pinhole}} = v \left(1 + k_1^y r^2 + k_2^y r^4 \right) ,$$

where $\begin{bmatrix} u, v \end{bmatrix}$ is the actual projection of a 3D point and $\begin{bmatrix} u_{\text{pinhole}}, v_{\text{pinhole}} \end{bmatrix}$ is the one the pinhole model predicts. In other words, the four parameters $\{k_1^x, k_2^x, k_1^y, k_2^y\}$ characterize the distortion. From now on, we will therefore write the full projection as

$$\mathbf{X} = \pi(\mathbf{x}) = f_d(f_p(\mathbf{x})), \qquad (5)$$
$$\mathbf{X} = \begin{bmatrix} x, y, z \end{bmatrix}, \qquad (5)$$
$$\mathbf{x} = \begin{bmatrix} u, v \end{bmatrix}, \qquad (5)$$

where f_p denotes the ideal pinhole projection of **Equation 3** and f_d the correction of **Equation 4**.

Triangulation. We can associate to each of the seven cameras a projection function π_c like the one in *Equation 5*, where *c* is the camera number. Given a 3D point and its projections \mathbf{x}_c in the images, its 3D coordinates can be estimated by minimizing the *reprojection error*

$$\underset{\mathbf{X}\in\mathbb{R}^{4}}{\operatorname{argmin}}\sum_{c=1}^{7}e_{c}\|\boldsymbol{\pi}_{c}(\mathbf{X})-\mathbf{x}_{c}\|_{2}^{2},$$
(6)

where e_c is one if the point was visible in image c and zero otherwise. In the absence of camera distortion, that is, when the projection π is a purely linear operation in homogeneous coordinates, this can be done for any number of cameras by solving a Singular Value Decomposition (SVD) problem (*Hartley and Zisserman, 2000*). In the presence of distortions, we replace the observed uand v coordinates of the projections by the corresponding $u_{pinhole}$ and $u_{pinhole}$ values of *Equation 5* before performing the SVD.

Camera calibration. Triangulating as described above requires knowing the projection matrices \mathbf{P}_c of *Equation 3* for each camera *c*, corresponding distortion parameters $\{k_1^x, k_2^x, k_1^y, k_2^y\}$ of *Equation 4*, together with the intrinsic parameters of focal length and principal point offset. In practice, we use the focal length and principal point offset provided by the manufacturer and estimate the remaining parameters automatically: the three translations and three rotations for each camera that define the corresponding matrix **M** of extrinsic parameters along with the distortion parameters.

To avoid having to design the exceedingly small calibration pattern that more traditional methods use to estimate these parameters, we use the fly itself as calibration pattern and minimize the reprojection error of *Equation 6* for all joints simultaneously while allowing the camera parameters to also change. In other words we look for

$$\underset{\substack{\pi_{c1 \leq c \leq 7} \\ \mathbf{X}_{j_{1 \leq j \leq n}}}{\operatorname{argmin}} \sum_{c=1}^{7} \sum_{j=1}^{m} e_{c,j} \rho(\pi_{c}(\mathbf{X}_{j}) - \mathbf{x}_{c,j}),$$

$$(7)$$

where X_j and $x_{c,j}$ are the 3D locations and 2D projections of the landmarks introduced above and ρ denotes the Huber loss. *Equation 7* is known as bundle-adjustment (*Hartley and Zisserman, 2000*).

332 Huber loss is defined as

$$\rho_{\delta}(a) = \begin{cases} \frac{1}{2}a^2 & \text{for } |a| \le \delta\\ \delta\left(|a| - \frac{1}{2}\delta\right) & \text{otherwise} \end{cases}$$

³³³ Replacing the squared loss by the Huber loss makes our approach more robust to erroneous ³³⁴ detections $\mathbf{x}_{c,j}$. We empirically set δ to 20 pixels. Note that we perform this minimization with respect ³³⁵ to ten degrees-of-freedom per camera: three translations, three rotations, and four distortions.

For this optimization to work properly, we need to initialize these ten parameters and we need to 336 reduce the number of outliers. To achieve this, the initial distortion parameters are set to zero. We 337 also produce initial estimates for the three rotation and three translation parameters by measuring 338 the distances between adjacent cameras and their relative orientations. To initialize the rotation 339 and translation vectors, we measure the distance and the angle between adjacent cameras, from 340 which we infer rough initial estimates. Finally, we rely on epipolar geometry (Hartley and Zisserman, 341 2000) to automate outlier rejection. Because the cameras form a rough circle and look inward, the 342 epipolar lines are close to being horizontal. Thus, corresponding 2D projections must belong to the 343 same image rows, or at most a few pixels higher or lower. In practice, this means checking if all 2D 344 predictions lie in nearly the same rows and discarding *a priori* those that do not. 345

346 **3D pose correction**

The triangulation procedure described above can produce erroneous results where the 2D estimates of landmarks are wrong. Additionally, it may result in implausible 3D poses for the entire animal because it treats each joint independently. To enforce more global geometric constraints, we rely bioRxiv preprint first posted online May. 20, 2019; doi: http://dx.doi.org/10.1101/640375. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a CC-Bpeep http://gob.2019.attional.license.

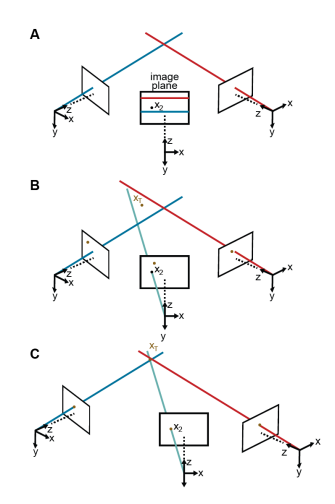


Figure 7. Camera calibration. **(A)** Correcting erroneous 2D pose estimations by using epipolar relationships. Only 2D pose estimates without large epipolar errors are used for calibration. x_2 represents a 2D pose estimate from the middle camera. Epipolar lines are indicated as blue and red lines on the image plane. **(B)** The triangulated point, X_T , uses the initial camera parameters. However, due to the coarse initialization of each camera's extrinsic properties, observations from each camera do not agree with one another and do not yield a reasonable 3D position estimate. **(C)** The camera locations are corrected, generating an accurate 3D position estimate by optimizing *Equation 7* using only the pruned 2D points.

on pictorial structures (*Felzenszwalb and Huttenlocher, 2005*) as described in *Figure 8*. Pictorial structures encode the relationship between a set of variables (in this case the 3D location of separate tracked points) in a probabilistic setting using a graphical model. This makes it possible to consider multiple 2D locations $\mathbf{x}_{c,j}$ for each landmark \mathbf{X}_c instead of only one. This increases the likelihood of finding the true 3D pose. **Generating multiple candidates.** Instead of selecting landmarks as the locations with the

Generating multiple candidates. Instead of selecting landmarks as the locations with the maximum probability in maps output by our Stacked Hourglass network, we generate multiple candidate 2D landmark locations $x_{c,j}$. From each probability map, we select ten local probability maxima that are at least one pixel apart from one another. Then, we generate 3D candidates by triangulating 2D candidates in every tuple of cameras. Because a single point is visible from at most four cameras, this results in at most $\binom{4}{2} \times 10^2$ candidates for each tracked point.

Choosing the best candidates. To identify the best subset of resulting 3D locations, we introduce the probability distribution $P(L|I, \theta)$ that assigns a probability to each solution L, consisting of 38 sets of 2D points observed from each camera. Our goal is then to find the most likely one. More formally, P represents the likelihood of a set of tracked points L, given the images, model bioRxiv preprint first posted online May. 20, 2019; doi: http://dx.doi.org/10.1101/640375. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a CC-By elocity above a license.

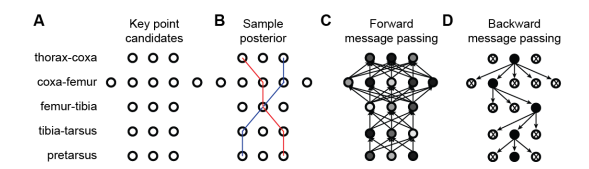


Figure 8. 3D pose correction for one leg using the MAP solution and pictorial structures. **(A)** Candidate 3D pose estimates for each keypoint are created by triangulating local maxima from probability maps generated by the Stacked Hourglass deep network. **(B)** For a selection of these candidate estimates, we can assign a probability using *Equation 8*. However, calculating this probability for each pair of points is computationally intractable. **(C)** By exploiting the chain structure of *Equation 8*, we can instead pass a probability distribution across layers using a belief propagation algorithm. Messages are passed between layers as a function of parent nodes, describing the belief of the child nodes on each parent node. Grayscale colors represent the calculated belief of each node where darker colors indicate higher belief. **(D)** Corrected pose estimates are obtained during the second backward iteration, by selecting the nodes with largest belief. We discard nodes (x's) that have non-maximal belief during backwards message passing. Note that beliefs have been adjusted after forward message passing.

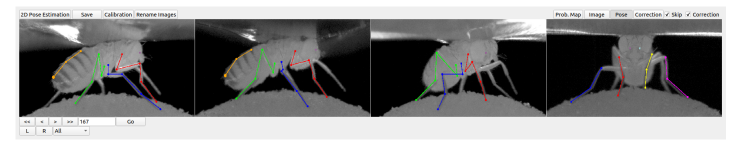


Figure 9. DeepFly3D graphical user interface (GUI). The top-left buttons enable operations like 2D pose estimation, camera calibration, and saving the final results. The top-right buttons can be used to visualize the data in different ways: as raw images, probability maps, 2D pose, or the corrected pose following pictorial structures. The bottom-left buttons permit frame-by-frame navigation. A full description of the GUI can be found in the online documentation: https://github.com/NeLy-EPFL/DeepFly3D

³⁶⁵ parameters, camera calibration, and geometric constraints. In our formulation, *I* denotes the seven

camera images $I = \{I_c\}_{1 \le c \le 7}$ and θ represents the set of projection functions π_c for camera *c* along with a set of length distributions $S_{i,j}$ between each pair of points *i* and *j* that are connected by a limb. *L* consists of a set of tracked points $\{L_i\}_{1 \le i \le n}$, where each L_i describes a set of 2D observations $I_{i,c}$ from multiple camera views. These are used to triangulate the corresponding 3D point locations $\overline{I_i}$. If the set of 2D observations is incomplete, as some points are totally occluded in some camera views, we triangulate the 3D point $\overline{I_i}$ using the available ones and replace the missing observations by projecting the recovered 3D positions into the images, $\pi_c(\overline{I_i})$ in **Equation 3**. In the end, we aim to

find the solution $\hat{L} = \operatorname{argmax}_{L} P(L|I, \theta)$. This is known as Maximum a Posteriori (MAP) estimation.

Using Bayes rule, we write

$$P(L|I,\theta) \propto P(I|L,\theta)P(L|\theta)$$
, (8)

where the two terms can be computed separately. We compute $P(I|J, \theta)$ using the probability maps $H_{j,c}$ generated by the Stacked Hourglass network for the tracked point j for camera c. For a single joint j seen by camera c, we model the likelihood of observing that particular point using $P(H_{j,c}|I_{j,c})$, which can be directly read from the probability maps as the pixel intensity. Ignoring the

dependency between the cameras, we write the overall likelihood as the product of the individual

380 likelihood terms

$$P(I|L,\theta) = P(H|L) \propto \prod_{i=1}^{n} \prod_{c=1}^{7} P(H_{j,c}|l_{i,c}) ,$$

³⁸¹ which can be read directly from the probability maps as pixel intensities and represent the network's

- ³⁸² confidence that a particular keypoint is located at a particular pixel. When a point is not visible from
- ³⁸³ a particular camera, we assume the probability map only contains a constant non-zero probability,
- which does not effect the final solution. We express $P(L|\theta)$ as

$$P(L|\theta) = P(L|\pi, S) = \prod_{(i,j)\in E} P\left(\bar{l}_i, \bar{l}_j|S_{i,j}\right) \prod_{j=1}^n \prod_{c=1}^7 e_{c,j} \|\pi_c(\bar{l}_j) - l_{c,j}\|_2^{-1},$$

where pairwise dependencies $P\left(\bar{l}_i, \bar{l}_j | S_{ij}\right)$ between two variables respect the segment length 385 constraint when the variables are connected by a limb. The length of segments defined by pairs of 386 connected 3D points follows a normal distribution. Specifically, we model $P(\bar{l}_i, \bar{l}_j | S_{i,j})$ as $S_{i,j}(\bar{l}_i, \bar{l}_j) =$ 387 $\mathcal{N}(\|\bar{l}_i - \bar{l}_i\| - \mu_{i,i}, \sigma_{i,i})$. We model the reprojection error for a particular point j as $\prod_{c=1}^7 e_{c,j} \|\pi_c(\bar{l}_j) - l_{c,j}\|_2^{-1}$ 388 which is set to zero using the variable $e_{c,i}$ denoting the visibility of the point j from camera c. If a 2D 380 observation for a particular camera is manually set by a user with the DeepFlv3D GUI, we take it to 390 be the only possible candidate for that particular image and we set P(L,|H) to 1, where i denotes 391 the manually assigned pixel location. 392

Solving the MAP problem using the Max-Sum algorithm. For general graphs, MAP estimation 393 with pairwise dependencies is NP-hard and therefore intractable. However, in the specific case of 394 non-cyclical graphs, it is possible to solve the inference problem using belief propagation (*Bishop*, 395 2006). Since the fly's skeleton has a root and contains no loops, we can use a message passing 396 approach (*Felzenszwalb and Huttenlocher, 2005*). It is closely related to Viterbi recurrence and 397 propagates the unary probabilities $P(L_i|L_i)$ between the edges of the graph starting from the root 398 and ending at the leaf nodes. This first propagation ends with the computation of the marginal 390 distribution for the leaf node variables. During the subsequent backward iteration, as $P(L_i)$ for 400 leaf node is computed, the point L_i with maximum posterior probability is selected in O(k) time, 401 where k is the upper bound on the number of proposals for a single tracked point. Next, the 402 distribution $P(L_i|L_i)$ is calculated, adjacent nodes for the leaf node. Continuing this process on 403 all of the remaining points results in a MAP solution for the overall distribution P(L), as shown in 404 **Figure 8.** with overall $O(k^2)$ computational complexity. 405

Learning the parameters. We learn the parameters for the set of pairwise distributions $S_{i,j}$ using a maximum likelihood process and assuming the distributions to be Gaussian. We model the segment length $S_{i,j}$ as the euclidean distance between the points \bar{l}_j and \bar{l}_j . We then solve for argmax_S $P(S|L, \theta)$, assuming segments have a Gaussian distribution resulting from the Gaussian noise in point observations L. This gives us the mean and variance, defining each distribution $S_{i,j}$. We exclude the same points that we removed from the calibration procedure, that exhibit high reprojection error.

In practice, we observe a large variance for pretarsus values. This is because occlusions occasionally shorten visible tarsal segments. To eliminate the resulting bias, we treat these limbs differently from the others and model the distribution of tibia-tarsus and tarsus-tip points as a Beta distribution, with parameters found using a similar Maximum Likelihood Estimator (MLE) formulation. Assuming the observation errors to be Gaussian and zero-centered, the bundle adjustment procedure can also be understood as an MLE of the calibration parameters (*Triggs et al., 2000*). Therefore, the entire set of parameters for the formulation can be learned using MLE. The pictorial structure formulation can be further expanded using temporal information, pe-

In protorial structure formulation can be further expanded using temporal information, pe nalizing large movements of a single tracked point between two consecutive frames. However, we
 abstained from using temporal information more extensively for several reasons. First, temporal
 dependencies would introduce loops in our pictorial structures, thus making exact inference NP hard as discussed above. This can be handled using loopy belief propagation algorithms (*Murphy*)

et al., 1999) but requires multiple message passing rounds, which prevents real-time inference without any theoretical guarantee of optimal inference. Second, the rapidity of *Drosophila* limb movements makes it hard to assign temporal constraints, even with fast video recording. Finally, we empirically observed that the current formulation, enforcing structured poses in a single temporal frame, already eliminates an overwhelming majority of false-positives inferred during the pose

430 estimation stage of the algorithm.

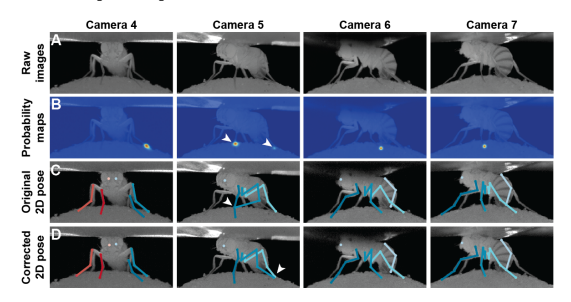


Figure 10. Pose correction using pictorial structures. **(A)** Raw input data from four cameras, focusing on the pretarsus of the middle left leg. **(B)** Probability maps for the pretarsus output from the Stacked Hourglass deep network. Two maxima (white arrowheads) are present on the probability maps for camera 5. The false-positive has a larger unary probability. **(C)** Raw predictions of 2D pose estimation without using pictorial structures. The pretarsus label is incorrectly applied (white arrowhead) in camera 5. By contrast, cameras 4, 6, and 7 are correctly labeled. **(D)** Corrected pose estimation using pictorial structures. The false-positive is removed due to the high error measured in *Equation 8*. The newly corrected pretarsus label for camera 5 is shown (white arrowhead).

Modifying DeepFly3D to study other animals. DeepFly3D can also be applied toward 3D pose estimation in other animals (e.g., humans (*Figure 11*), primates, rodents, and other insects). Importantly, DeepFly3D does not assume a circular camera arrangement, or that there is one degree of freedom in the camera network. We illustrate this flexibility by using DeepFly3D to analyze the Human 3.6M Dataset (http://vision.imar.ro/human3.6m/description.php) generated from four synchronized cameras (*Ionescu et al., 2014*).

Generally, for any new dataset, the user first needs to provide an initial set of manual annotations. 437 Then, in *skeleton.py*, the user should describe the number of tracked points and their relationships 438 to one another (e.g., are they connected). Then, in Config.py, the user should set the number of 439 cameras, and the resolutions of input images and output probability maps. DeepFly3D will then use 440 these initial manual annotations to (i) train the 2D Stacked Hourglass network. (ii) perform camera 441 calibration without an external checkerboard pattern, (iii) learn the epipolar geometry to perform 442 outlier detection, and (iv) learn the segment length distributions $S_{i,i}$. After this initial bootstrapping, 443 DeepFly3D can be then used with pictorial structures and active learning to iteratively improve pose 444 estimation accuracy. 445 The initial manual annotations can be performed using the DeepFly3D Annotation GUI. After-446

wards, these annotations can be downloaded from the Annotation GUI as a CSV file using the *Save* button (*Figure 6*). Once the CSV file is placed in the images folder, DeepFly3D will automatically
 read and display the annotations. To train the Stacked Hourglass network, use the *csv-path* flag
 while running *pose2d.py* (found in *deepfly/pose2d*). DeepFly3D will then train the Stacked Hourglass

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Figure 11. DeepFly3D graphical user interface (GUI) used with the Human3.6M dataset *Ionescu et al. (2014*). To use the DeepFly3D GUI on any new dataset (*Drosophila* or otherwise), users can provide an initial small set of manual annotations. Using these annotations, the software calculates the epipolar geometry, performs camera calibration, and trains the 2D pose estimation deep network. A description of how to adopt DeepFly3D for new datasets can be found in the Methods section and, in greater detail, online: https://github.com/NeLy-EPFL/DeepFly3D

- network by performing transfer learning using the large MPII dataset and the smaller set of user manual annotations.
- 453 To perform camera calibration, the user should select the *Calibration* button on the GUI *Figure 9*.
- ⁴⁵⁴ DeepFly3D will then perform bundle adjustment (*Equation 7*) and save the camera parameters
- in *calibration.pickle* (found in the images folder). The path of this file should then be added to
- 456 Config.py to initialize calibration. These initial calibration parameters will then be used in further
- ⁴⁵⁷ experiments for fast and accurate convergence. If the number of annotations is insufficient for
- accurate calibration, or if bundle adjustment is converging too slowly, an initial rough estimate of the camera locations can be set in *Config.pv*. As long as a calibration is set in *Config.pv*. DeepFlv3D
- the camera locations can be set in *Config.py*. As long as a calibration is set in *Config.py*, DeepFly3D
 will use it as a projection matrix to calculate the epipolar geometry between cameras. This step is
- ⁴⁶¹ necessary to perform outlier detection on further calibration operations.
- ⁴⁶² DeepFly3D will also learn the distribution $S_{i,j}$, whose non-zero entries are found in *skeleton.py*. ⁴⁶³ One can easily calculate these segment length distribution parameters using the functions provided ⁴⁶⁴ with DeepFly3D. *CameraNetwork* class (found under deepfly/GUI/), will then automatically load the ⁴⁶⁵ points and calibration parameters from the images folder. The function *CameraNetwork.triangulate* ⁴⁶⁶ will convert 2D annotation points into 3D points using the calibration parameters. The $S_{i,j}$ pa-⁴⁶⁷ rameters can then be saved using the *pickle* library (the save path can be set in *Config.py*). The ⁴⁶⁸ *calcBoneParams* method will then output the segment lengths' mean and variance. These values
- will then be used with pictorial structures (*Equation 8*).
- We provide further technical details for how to adapt DeepFly3D to other multi-view datasets online ¹.

472 Experimental setup

We positioned seven Basler acA1920-155um cameras (FUIIFILM AG. Niederhaslistrasse, Switzerland) 473 94 mm away from the tethered fly, resulting in a circular camera network with the animal in the 474 center (*Figure 12*). We acquired 960×480 pixel video data at 100 FPS under 850 nm infrared ring 475 light illumination (Stemmer Imaging, Pfäffikon Switzerland). Cameras were mounted with 94 mm 476 W.D. / 1.00x InfiniStix lenses (Infinity Photo-Optical GmbH, Göttingen). Optogenetic stimulation LED 477 light was filtered out using 700 nm longpass optical filters (Edmund Optics, York UK). Each camera's 478 depth of field was increased using 5.8 mm aperture retainers (Infinity Photo-Optical GmbH). To 479 automate the timing of optogenetic LED stimulation and camera acquisition triggering, we use an 480

¹https://github.com/NeLy-EPFL/DeepFly3D

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481 Arduino (Arduino, Sommerville MA USA) and custom software written using the Basler camera API.

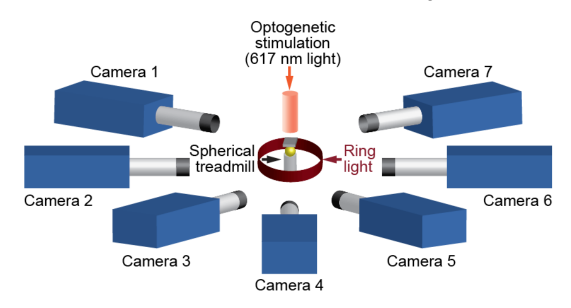


Figure 12. A schematic of the seven camera spherical treadmill and optogenetic stimulation system that was used in this study.

Drosophila transgenic lines. UAS-CsChrimson (Klapoetke et al., 2014) animals were obtained
 from the Bloomington Stock Center (Stock #55135). MDN-1-Gal4 (Bidaye et al., 2014) (VT44845 DBD; VT50660-AD) was provided by B. Dickson (Janelia Research Campus, Ashburn USA). aDN-Gal4
 (Hampel et al., 2015)(R76F12-AD; R18C11-DBD), was provided by J. Simpson (University of California,
 Santa Barbara USA). Wild-type, PR animals were provided by M. Dickinson (California Institute of
 Technology, Pasadena USA).

Optogenetic stimulation experiments. Experiments were performed in the late morning or 488 early afternoon Zeitgeber time (Z.T.), inside a dark imaging chamber. An adult female animal 2-3 489 days-post-eclosion (dpe), was mounted onto a custom stage (Chen et al., 2018) and allowed to 490 acclimate for 5 minutes on an air-supported spherical treadmill (*Chen et al., 2018*). Optogenetic 491 stimulation was performed using a 617 nm LED (Thorlabs, Newton, NJ USA) pointed at the dorsal 492 thorax through a hole in the stage, and focused with a lens (LA1951, 01" f = 25.4 mm, Thorlabs, 493 Newton, NI USA). Tethered flies were otherwise allowed to behave spontaneously. Data were 494 acquired in 9 s epochs: 2 s baseline, 5 s with optogenetic illumination, and 2 s without stimulation. 495 Individual flies were recorded for 5 trials each, with one-minute intervals. Data were excluded 496 from analysis if flies pushed their abdomens onto the spherical treadmill—interfering with limb 497 movements—or if flies struggled during optogenetic stimulation, pushing their forelimbs onto the 498 stage for prolonged periods of time. 499

500 Unsupervised behavioral classification

To create unsupervised embeddings of behavioral data, we mostly followed the approach taken 501 by (Todd et al., 2017; Berman et al., 2014). We smoothed 3D pose traces using a 1€ Filter. Then 502 we converted them into angles to achieve scale and translational invariance (Casiez et al., 2012). 503 Angles were calculated by taking the dot product from sets of three connected 3D positions. For 504 the antenna, we calculated the angle of the line defined by two antennal points with respect to the 505 ground-plane. This way, we generated four angles per leg (two body-coxa, one coxa-femur, and 506 one femur-tibia), two angles for the abdomen (top and bottom abdominal stripes), and a single 507 angle for the antennae (head tilt with respect to the axis of gravity). In total, we obtained a set of 34 508 angles, extracted from 38 3D points. 509

We transformed angular time series using a Continous Wavelet Transform (CWT) to create a 510 posture-dynamics space. We used the Morlet Wavelet as the mother wavelet, given its suitability to 511 isolate periodic chirps of motion. We chose 25 wavelet scales to match dyadically spaced center 512 frequencies between 5Hz and 50Hz. Then, we calculate spectrograms for each postural time-series 513 by taking the magnitudes of the wavelet coefficients. This yields a $34 \times 25 = 850$ -dimensional 514 time-series, which was then normalized over all frequency channels to unit length, at each time 515 instance. Then, we could treat each feature vector from each time instance as a distribution over all 516 frequency channels. 517

Later, from the posture-dynamics space, we computed a two-dimensional representation of 518 behavior by using the non-linear embedding algorithm, t-SNE Magten and Hinton (2008), t-SNE em-519 bedded our high-dimensional posture-dynamics space onto a 2D plane, while preserving the high-520 dimensional local structure, while sacrificing larger scale accuracy. We used the Kullback-Leibler 521 (KL) divergence as the distance function in our t-SNE algorithm. KL assesses the difference between 522 the shapes of two distributions, justifying the normalization step in the preceding step. By analyzing 523 a multitude of plots generated with different perplexity values, we empirically found perplexity 35 524 to best suit the features of our posture-dynamics space. 525

From this generated discrete space, we created a continuous 2D distribution, that we could then segment into behavioral clusters. We started by normalizing the 2D t-SNE projected space into a 1000×1000 matrix. Then, we applied a 2D Gaussian convolution with a kernel of size $\sigma = 10$ px. Finally, we segmented this space by inverting it and applying a Watershed algorithm that separated adjacent basins, yielding a behavioral map.

531 Author Contributions

532 SG - Conceptualization, Methodology, Software, Validation, Formal Analysis, Investigation, Data

- 533 Curation, Writing Original Draft Preparation, Writing Review & Editing, Visualization
- 534

535 HR - Conceptualization, Methodology, Software, Formal Analysis, Writing - Original Draft Preparation,

536 Writing - Review & Editing, Supervision, Project Administration

- 537
- DM Investigation, Data Curation, Writing Review & Editing
- 538 539

JC - Software, Data Curation, Writing - Review & Editing

541

544

PR - Conceptualization, Methodology, Resources, Writing - Original Draft Preparation, Writing -

Review & Editing, Supervision, Project Administration, Funding Acquisition

PF - Conceptualization, Methodology, Resources, Writing - Review & Editing, Supervision, Project
 Administration, Funding Acquisition

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sso support from an EPFL SV iPhD grant

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- **554 Competing interests**
- 555 The authors declare that no competing interests exist.

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