

Objective comparison of particle tracking methods

Nicolas Chenouard^{1–3,25}, Ihor Smal^{4,5,25}, Fabrice de Chaumont^{1,25}, Martin Maška^{6,7,25}, Ivo F Sbalzarini⁸, Yuanhao Gong⁸, Janick Cardinale⁸, Craig Carthel⁹, Stefano Coraluppi⁹, Mark Winter¹⁰, Andrew R Cohen¹⁰, William J Godinez^{11,12}, Karl Rohr^{11,12}, Yannis Kalaidzidis^{13,14}, Liang Liang¹⁵, James Duncan¹⁵, Hongying Shen¹⁶, Yingke Xu¹⁷, Klas E G Magnusson¹⁸, Joakim Jaldén¹⁸, Helen M Blau¹⁹, Perrine Paul-Gilloteaux²⁰, Philippe Roudot²¹, Charles Kervrann²¹, François Waharte²⁰, Jean-Yves Tinevez²², Spencer L Shorte²², Joost Willemsse²³, Katherine Celler²³, Gilles P van Wezel²³, Han-Wei Dan²⁴, Yuh-Show Tsai²⁴, Carlos Ortiz de Solórzano⁶, Jean-Christophe Olivo-Marin^{1,26} & Erik Meijering^{4,5,26}

Particle tracking is of key importance for quantitative analysis of intracellular dynamic processes from time-lapse microscopy image data. Because manually detecting and following large numbers of individual particles is not feasible, automated computational methods have been developed for these tasks by many groups. Aiming to perform an objective comparison of methods, we gathered the community and organized an open competition in which participating teams applied their own methods independently to a commonly defined data set including diverse scenarios. Performance was assessed using commonly defined measures. Although no single method performed best across all scenarios, the results revealed clear differences between the various approaches, leading to notable practical conclusions for users and developers.

Technological developments in the past two decades have greatly advanced the field of bioimaging and have enabled the investigation of dynamic processes in living cells at unprecedented spatial and temporal resolution. Examples include the study of cell membrane dynamics¹, cytoskeletal filaments², focal adhesions³, viral infection⁴, intracellular transport⁵, gene transcription⁶ and genome maintenance⁷. Apart from state-of-the-art light microscopy^{8,9} and fluorescent labeling^{10,11}, a key technology in the quest for quantitative analysis of intracellular dynamic

processes is particle tracking. Here, a ‘particle’ may be anything from a single molecule to a macromolecular complex, organelle, virus or microsphere¹², and the task of detecting and following individual particles in a time series of images is often (somewhat confusingly) referred to as ‘single-particle tracking’. As the number of particles may be very large (hundreds to thousands), requiring ‘multiple-particle tracking’^{13–15}, manual annotation of the image data is not feasible, and computer algorithms are needed to perform the task.

At present, dozens of software tools are available for particle tracking¹⁶. The image analysis methods on which they are based can generally be divided into two steps: (i) particle detection (the spatial aspect), in which spots that stand out from the background according to certain criteria are identified and their coordinates estimated in every frame of the image sequence, and (ii) particle linking (the temporal aspect), in which detected particles are connected from frame to frame using another set of criteria to form tracks. The two steps are commonly performed only once, but they may also be applied iteratively. For each of these steps, many methods have been devised over the years^{17–22}, often originating from other areas of data analysis^{23,24}. With so many methods currently known, the question arises as to what distinguishes them and how they perform relative to one another under different experimental conditions.

¹Institut Pasteur, Unité d'Analyse d'Images Quantitative, Centre National de la Recherche Scientifique Unité de Recherche Associée 2582, Paris, France. ²Biomedical Imaging Group, École Polytechnique Fédérale de Lausanne, Lausanne, Switzerland. ³New York University Neuroscience Institute, New York University Medical Center, New York, New York, USA. ⁴Department of Medical Informatics, Erasmus University Medical Center, Rotterdam, The Netherlands. ⁵Department of Radiology, Erasmus University Medical Center, Rotterdam, The Netherlands. ⁶Center for Applied Medical Research, University of Navarra, Pamplona, Spain. ⁷Centre for Biomedical Image Analysis, Masaryk University, Brno, Czech Republic. ⁸MOSAIC Group, Max Planck Institute of Molecular Cell Biology and Genetics, Dresden, Germany. ⁹CompuNetix Inc., Monroeville, Pennsylvania, USA. ¹⁰Department of Electrical and Computer Engineering, Drexel University, Philadelphia, Pennsylvania, USA. ¹¹Department of Bioinformatics and Functional Genomics, Institute of Pharmacy and Molecular Biotechnology, University of Heidelberg, Heidelberg, Germany. ¹²Division of Theoretical Bioinformatics, German Cancer Research Center, Heidelberg, Germany. ¹³Max Planck Institute of Molecular Cell Biology and Genetics, Dresden, Germany. ¹⁴Belozersky Institute of Physico-Chemical Biology, Moscow State University, Moscow, Russia. ¹⁵Department of Electrical Engineering, Yale University, New Haven, Connecticut, USA. ¹⁶Department of Cell Biology, Yale University, New Haven, Connecticut, USA. ¹⁷Department of Biomedical Engineering, Zhejiang University, Hangzhou, China. ¹⁸Department of Signal Processing, ACCESS Linnaeus Centre, KTH Royal Institute of Technology, Stockholm, Sweden. ¹⁹Baxter Laboratory for Stem Cell Biology, Department of Microbiology and Immunology, Stanford University School of Medicine, Stanford, California, USA. ²⁰Cell and Tissue Imaging Facility, Institut Curie, Paris, France. ²¹Inria Rennes, Bretagne Atlantique, Rennes, France. ²²Plateforme d'Imagerie Dynamique, Imagopole, Institut Pasteur, Paris, France. ²³Molecular Biotechnology Group, Institute of Biology, Leiden University, Leiden, The Netherlands. ²⁴Department of Biomedical Engineering, Chung Yuan Christian University, Chung Li City, Taiwan, China. ²⁵These authors contributed equally to this work. ²⁶These authors jointly directed this work. Correspondence should be addressed to: J.-C.O.-M. (jcolivo@pasteur.fr) or E.M. (meijering@imagescience.org).

Several comparison studies have been published in recent years. Cheezum *et al.* compared four basic methods for localization of a single particle that are often used for particle tracking and concluded that Gaussian fitting performs best by several criteria²⁵. A follow-up study, refining the conclusions by evaluating various practical aspects, was presented by Carter *et al.*²⁶. A more extensive study, evaluating nine methods (including two machine learning methods) for multiple-particle detection, was conducted by Smal *et al.*²⁷. They concluded that all methods perform well for sufficiently high signal-to-noise ratio (SNR ≥ 5); however, for low-quality images, learning-based methods are slightly superior, although other methods may yield comparable results and are easier to use. A similar study, published by Ruusuvaari *et al.*²⁸, rightly added that “algorithms should be chosen with care.” Finally, Godinez *et al.* compared eight different methods for tracking virus particles and found probabilistic methods to be superior²⁹.

Though interesting, the cited studies were limited to either one aspect of the task (detection rather than tracking) or one application (tracking of viruses rather than a broader set of particles). Moreover, the methods were implemented by the same group who performed the evaluation rather than by the original inventors. Obtaining a more complete picture of performance by combining the results of independent studies is usually hampered by their being based on different data sets and different evaluation criteria. Such fundamental problems have been recognized in the field of medical image analysis for more than 5 years and have resulted in the organization of international competitions (see <http://www.grand-challenge.org/>). The rationale behind such competitions is that the most objective evaluation of methods is achieved by having research groups apply their own methods independently, on a commonly defined data set and using commonly defined evaluation criteria. The first study in this spirit to be organized in the field of bioimage analysis was the digital reconstruction of axonal and dendritic morphology (DIADEM) challenge³⁰. For particle tracking, the organization of a competition was first advocated by Saxton¹² and in an editorial³¹.

Here we present an objective comparison of particle tracking methods based on an open competition that we organized in 2012 (see <http://www.bioimageanalysis.org/track/>). By announcements made through various media (at conferences, on the Web and via mailing lists and targeted emails) over 2 months, research groups worldwide were invited to participate. Next, registered teams were given 1 month to prepare their methods using representative training data and corresponding ground truth provided on the website. After release of the actual competition data, without ground truth, the teams were given 3 weeks to submit tracking results to an independent evaluator (one member of the organizing team who was not a contestant and the only one to have the ground truth). Preliminary results were presented and discussed at a workshop organized at the 2012 IEEE International Symposium on Biomedical Imaging. All participating teams sent their software to the independent evaluator who verified the results and performed an objective measurement of the computation times needed by the competing methods. A full analysis of the results and a discussion of the practical conclusions of our study are presented in this paper.

RESULTS

Participating teams and methods

A total of 14 teams (**Table 1**) took up the challenge and submitted tracking results. Together they used many different methods^{32–57} (**Table 1** and **Supplementary Note 1**) based on well-known as well as newly developed concepts. Approaches to particle detection ranged from simple thresholding or local-maxima finding to morphological processing, linear filtering (in particular, Gaussian, Laplacian of Gaussian and difference of Gaussian), linear and nonlinear model fitting, and centroid estimation schemes. Most detection methods were based on a combination of two or more of these. Approaches to linking of detected particles ranged from simple nearest-neighbor to multiframe association, including multiple hypothesis tracking, dynamic programming and combinatorial schemes, with or without explicit use of motion models and state estimation (Kalman filtering). Each tracking method consisted of a specific combination of detection and linking approaches as deemed appropriate by the corresponding team, who also determined suitable parameter settings for their method (**Supplementary Table 1**).

Data sets and ground truth

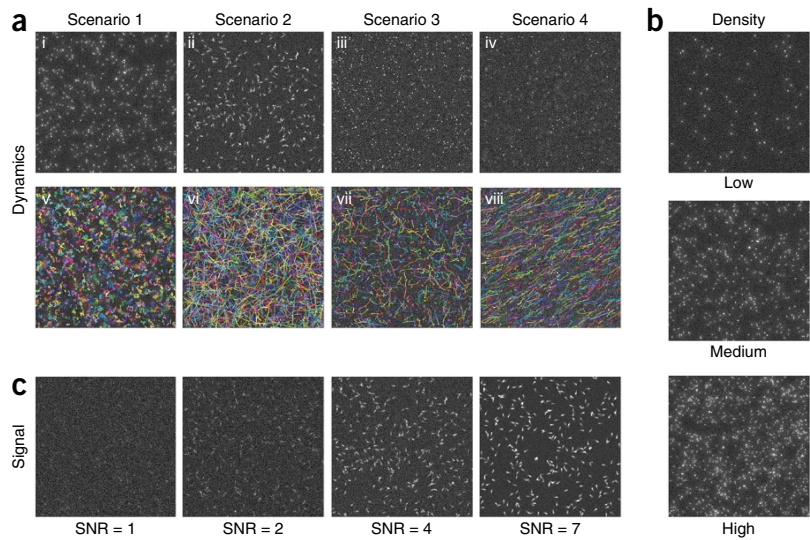
To allow an objective, quantitative comparison of the methods for a range of practical conditions, representative image data with exact ground truth was needed. Generally, ground truth is not available for real image data, and manual annotation by human observers is subjective, labor intensive for large numbers of particles and known^{14,19,58} to be potentially inferior to computational tracking in the first place, leading to inappropriate reference data. Therefore, we chose to simulate image data for this study (**Fig. 1**, **Table 2** and **Supplementary Videos 1–10**). We identified three main factors affecting tracking performance in practice (**Supplementary Note 2**): dynamics (type of motion), density (number of particles within the fixed field of view) and signal (relative to noise). For particle dynamics we considered four types of motion representative of a variety of biological scenarios, namely Brownian (random-walk) motion similar to that of vesicles in the cytoplasm, directed (near constant-velocity) motion such as microtubule transport and random switching between these two motion models, with either random or constrained orientation for the directed component, as with membrane receptors or infecting viruses, respectively (**Fig. 1a**, **Table 2** and **Supplementary Videos 1–4** and **10**). For particle density we considered three levels (**Fig. 1b** and **Supplementary Videos 5**, **1** and **6**, respectively): low (~100 particles), medium (~500 particles) and high (~1,000 particles), with random appearance and disappearance of particles. For particle signal relative to the noise, we considered four levels (**Fig. 1c** and **Supplementary Videos 7**, **8**, **2** and **9**, respectively): SNR = 1, 2, 4 and 7, where SNR = 4 was known from previous studies^{25,27} to be a critical level. Here, $SNR = (I_o - I_b) / \sqrt{I_o}$, with I_o denoting the peak object (particle) intensity and I_b the mean background intensity. Together this resulted in 48 cases. In all cases we modeled particles as labeled with GFP and imaged with fluorescence microscopy in either wide-field or confocal mode. The exact number of particles in any frame of a simulated time series, and the initiation, termination and displacement of particles from frame to frame, was governed by realistic random processes. The resulting data contained ambiguities similar to those in real data, including noise,

Table 1 | Participating teams and tracking methods

Method	Authors	Detection				Linking				Dim.	Refs.
		Prefilter	Approaches	Remarks	Principle	Approaches	Remarks	Dim.			
1	I.F. Sbalzarini Y. Gong J. Cardinale C. Carthel S. Coraluppi	–	M, C	Iterative intensity-weighted centroid calculation	Combinatorial optimization	MF, MT, GC	Greedy hill-climbing optimization with topological constraints	2D & 3D	32		
2	N. Chenouard F. de Chaumont J.-C. Olivo-Marin	Disk	M, T	Adaptive local-maxima selection	Multiple hypothesis tracking	MF, MT, MM	Motion models are user specified (near-constant position and/or velocity)	2D & 3D	33,34		
3	M. Winter A.R. Cohen	Wavelets	M, T	Maxima after thresholding two-scale wavelet products	Multiple hypothesis tracking	MF, MT, MM, GC	Motion models are user specified (near-constant position and/or velocity)	2D & 3D	35–37		
4	M. Winter A.R. Cohen	Gaussian, median and morphology	M, T, C	Adaptive Otsu thresholding	Multitemporal association tracking	MF, MT, GC	Post-tracking refinement of detections	2D & 3D	38,39		
5	W.J. Godinez K. Rohr	Laplacian of Gaussian or Gaussian fitting	M, T, F, C	Either thresholding + centroid or maxima + Gaussian fitting	Kalman filtering + probabilistic data association	MF, MM	Interacting multiple models using motion models as specified	2D & 3D	29,40		
6	Y. Kalaidzidis	Windowed floating mean background subtraction	T, F	Lorentzian function fitting to structures above noise level	Dynamic programming	MF, GC	Track assignment by the weighted sum of multiple features	2D	41		
7	L. Liang J. Duncan H. Shen Y. Xu	Laplacian of Gaussian	M, T, F	Gaussian mixture model fitting	Multiple hypothesis tracking	MF, MM	Interacting multiple models with forward and backward linking	2D	42		
8	K.E.G. Magnusson J. Jaldén H.M. Blau	Deconvolution	M, T, F	Watershed-based clump splitting and parabola fitting	Viterbi algorithm on state-space representation	MF, MT	Brownian motion is assumed in all cases	2D & 3D	43,44		
9	P. Paul-Giloteaux	Laplacian of Gaussian or Gaussian filtering	M, T, F	Either maxima with pixel precision (2D) or thresholding + Gaussian fitting (3D)	Nearest neighbor + global optimization	MF, MT, GC	Global optimization of associations using simulated annealing	2D & 3D	45,46		
10	P. Roudot C. Kervrann F. Waharte	Structure tensor	T, F	Histogram-based thresholding and Gaussian fitting	Gaussian template matching	–	Only local and per-trajectory particle linking	2D	47–49		
11	I. Smal E. Meijering	Wavelets	M, F, C	Gaussian fitting (round particles) or centroid calculation (elongated particles)	Sequential multiframe assignment	MF, MT, MM, GC	Global linking cost minimization	2D	35,50,51		
12	J.-Y. Tinevez S.L. Shorte	Difference of Gaussian	M, T, F	Parabolic fitting to localized maxima	Linear assignment problem	MT, GC	Two-step approach (frame-to-frame and segment linking)	2D & 3D	52,53		
13	J. Willemsse K. Celler G.P. van Wezel	Gaussian and top hat	T, C	Watershed-based clump splitting	Nearest neighbor	MM, GC	Allows merging and splitting of particles and uses a linear motion model	2D & 3D	54,55		
14	H.-W. Dan Y.-S. Tsai	Gaussian, Wiener and top hat	T, C	Morphological opening-based clump splitting	Nearest neighbor + Kalman filtering	MM	Essentially a 2D method keeping track of maximum intensity in z	2D & 3D	56,57		

See **Supplementary Note 1** for further details on methods 1–14. Dim., dimensionality. Detection approaches: M, maxima detection; T, thresholding; F, fitting; C, centroid estimation. Linking approaches: MF, multiframe; MT, multitrack; MM, motion models; GC, gap closing.

Figure 1 | Simulated image data. Representative images of the three main factors (particle dynamics, density and signal) affecting tracking performance are shown. **(a)** Four biological scenarios were simulated, of which we show snapshot images (**i–iv**) and trajectories (**v–viii**) in arbitrary colors: particles showing random-walk motion imaged in two dimensions over time (2D+time) using wide-field microscopy (**i,v**); larger (elongated) particles represented by asymmetric Gaussians showing directed motion in 2D+time (**ii,vi**); particles switching between random-walk and randomly oriented directed motion imaged in 2D+time using confocal microscopy (**iii,vii**); and particles switching between random-walk and directed motion with restricted orientation imaged in 3D+time (only one slice is shown) using confocal microscopy (**iv,viii**). **(b,c)** Three density levels (**b**; low, medium and high) and four SNR levels (**c**; 1, 2, 4 and 7) were simulated.



clutter, visual merging and splitting, and intersecting and parallel trajectories. In both the training and the competition phase of the study, participants were given only limited information about the data (Table 2).

Quantitative performance measures

A key problem when evaluating any method for tracking large numbers of particles is how to optimally pair the set of estimated tracks, Y , with the set of ground-truth tracks, X , which are likely to contain different numbers of elements (tracks and points within tracks). To solve this, we extended Y with dummy tracks and applied optimal subpattern assignment using the Munkres algorithm⁵⁹, which yielded the globally best possible pairing (minimal total distance) of each ground-truth track, θ_k^X , with either an estimated track (if available) or a dummy track (in the absence of a suitable estimated track), θ_k^Z , where Z denotes the dummy-extended and ordered version of Y . In the pairing process, the distance $d(\theta_k^X, \theta_k^Z)$ between two tracks was computed as the sum, over all time points t of the image sequence, of the gated Euclidean distance between the corresponding track points, $d(\theta_k^X, \theta_k^Z) = \sum_t |\theta_k^X(t) - \theta_k^Z(t)|_{2,\varepsilon}$, with $\theta(t)$ denoting the spatial position of a track at time t and $|\cdot|_{2,\varepsilon} = \min(|\cdot|_2, \varepsilon)$. If at any t a track point was missing, a dummy point was taken. The gate ε served both to determine whether the points of paired tracks were matching at any t and to apply a fixed penalty to non-matching points. In this study, ε was set to 5 pixels, which was on the order of the Rayleigh distance in our data (Supplementary Note 2). The total distance $d(X, Y)$ between track sets X and Y , minimized by the Munkres algorithm by optimizing Z , was simply the sum over all k of the distances $d(\theta_k^X, \theta_k^Z)$ between paired tracks.

On this basis, we considered 14 different aspects of tracking accuracy, which we summarized into five performance measures. The five measures (Supplementary Note 3) were as follows.

1. $\alpha(X, Y) = 1 - d(X, Y) / d(X, \emptyset)$. \emptyset denotes a set of dummy tracks; hence, $d(X, \emptyset)$ is the maximum possible total distance (error) from the ground truth. The measure ranges from 0 (worst) to 1 (best), indicating the overall degree of matching of ground-truth and estimated tracks without taking into account spurious (nonpaired estimated) tracks.

2. $\beta(X, Y) = (d(X, \emptyset) - d(X, Y)) / (d(X, \emptyset) + d(\bar{Y}, \emptyset))$. \bar{Y} denotes the set of spurious tracks, and $d(\bar{Y}, \emptyset)$ is the corresponding penalty term. The measure ranges from 0 (worst) to α (best) and is essentially α with a penalization of nonpaired estimated tracks.

3. $JSC = TP / (TP + FN + FP)$. This is the Jaccard similarity coefficient for track points. It ranges from 0 (worst) to 1 (best) and characterizes overall particle detection performance. TP (true positives) denotes the number of matching points in the optimally paired tracks; FN (false negatives), the number of dummy points in the optimally paired tracks; and FP (false positives), the number of nonmatching points including those of the spurious tracks.

4. $JSC_\theta = TP_\theta / (TP_\theta + FN_\theta + FP_\theta)$. This is the Jaccard similarity coefficient for entire tracks instead of single track points. Similarly to JSC, it ranges from 0 (worst) to 1 (best). TP_θ denotes the number of estimated tracks paired with ground-truth tracks; FN_θ , the number of dummy tracks paired with ground-truth tracks; and FP_θ , the number of spurious tracks.

5. RMSE, the r.m.s. error, indicates the overall localization accuracy of matching points in the optimally paired tracks (the TP as in JSC).

Table 2 | Basic properties of the image data

Parameter	Scenario 1	Scenario 2	Scenario 3	Scenario 4
Nickname	Vesicles	Microtubules	Receptors	Viruses
Dynamics	Brownian	Directed	Switching	Switching
PSF model	Wide field	Nonisotropic Gaussian	Confocal	Confocal
Dimensionality	2D+time	2D+time	2D+time	3D+time
Image size (pixels)	512 × 512	512 × 512	512 × 512	512 × 512
Stack size (slices)	1	1	1	10
Length (frames)	100	100	100	100
Density ^a (low, medium, high)	100, 500, 1,000	60, 400, 700	100, 500, 1,000	100, 500, 1,000
SNR (levels)	1, 2, 4, 7	1, 2, 4, 7	1, 2, 4, 7	1, 2, 4, 7
Intersection fraction ^b (%)	0.7, 3.4, 6.8	0.8, 4.9, 8.8	0.5, 2.6, 5.4	0.3, 1.2, 2.4

^aOn average per time point. ^bFor the low-, medium- and high-density data per scenario, using the Rayleigh distance as the criterion to determine intersection, and averaged over the different SNR levels.

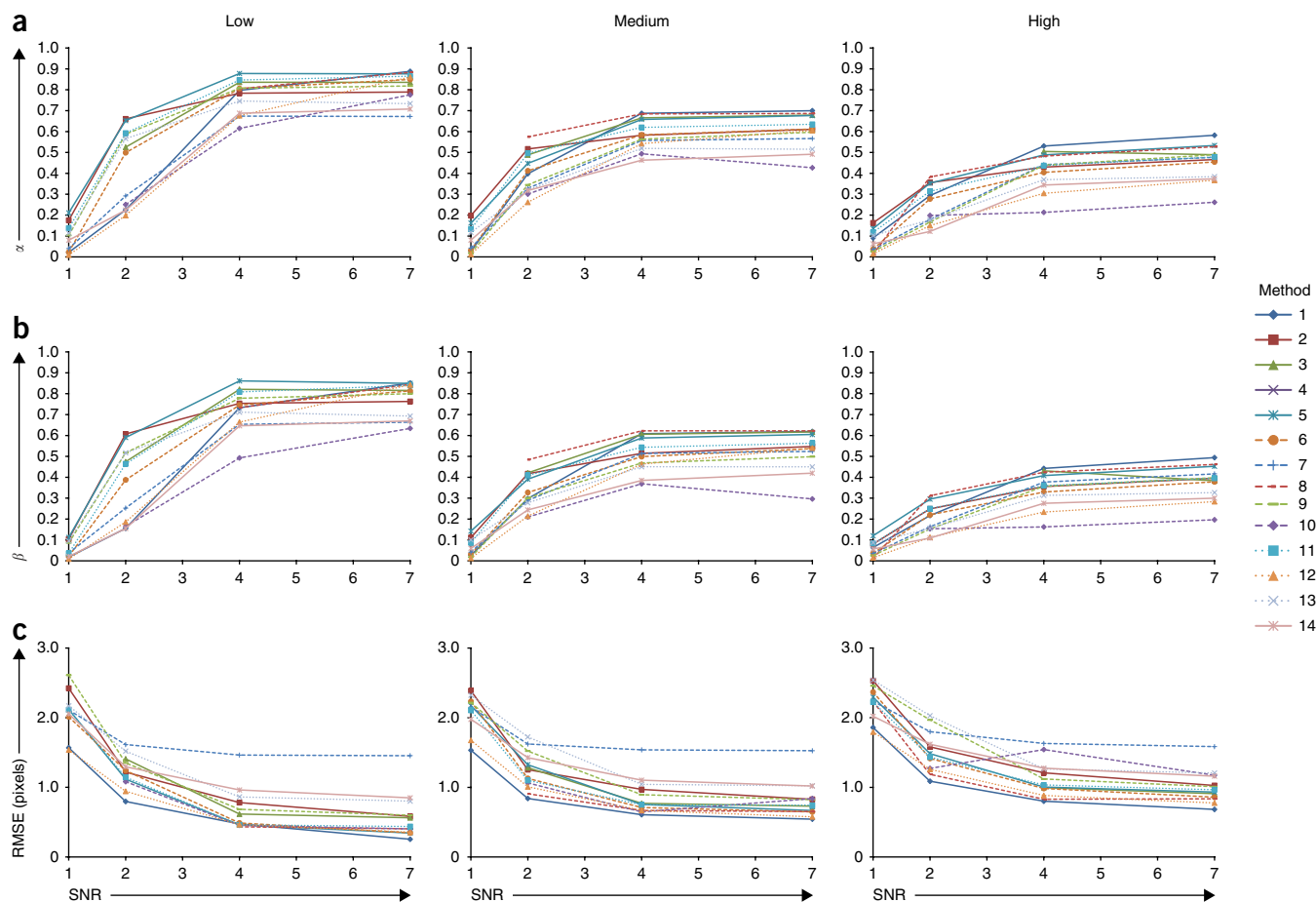


Figure 2 | Sample performance results. Values of three performance measures (α , β and RMSE) are plotted as a function of density (low, medium and high) and SNR for scenario 1. (a) α values (scoring the match between ground-truth and estimated tracks) for each density. (b) β values (α values with a penalty for nonmatching estimated tracks) for each density. (c) RMSE values (scoring localization accuracy) for each density. For some methods, the lines are incomplete, indicating missing (not submitted) tracking results.

Submission of tracking results

Not all teams submitted results for all 48 cases. Some of their methods were not designed to deal with severe noise or more complex shapes or dynamics. Some methods (Table 1) were developed for tracking in only two-dimensional (2D) time series and could not be applied to the 3D cases. And some teams reported insufficient time to complete the tracking of all cases within the 3-week competition phase. Nevertheless, of the 48 (data) \times 14 (teams) = 672 possible tracking results, 505 (75%) were submitted to the independent evaluator, who computed the values of all performance measures (Supplementary Table 2), verified the results (Supplementary Table 3) and measured the computation times needed by the methods (Supplementary Table 4).

Performance of the methods

For each tracking method, the values of the performance measures were computed for each data case for which tracking results were submitted. Basing our analysis on the computed values, we studied the performance of the different methods as a function of particle dynamics (the different biological scenarios modeled), density and signal level (Fig. 2 and Supplementary Table 2) as well as in terms of their required computation times (Supplementary Table 4). We subsequently ranked the methods according to best performance per case (Supplementary Tables 2 and 4)

as described in the Online Methods. From these rankings we considered the top 3 best-performing methods (Fig. 3) and studied the effects of decreasing the value of the gate parameter ϵ (Supplementary Table 5).

The global observation from the results is that no one particle tracking method performed best for all data. Nevertheless, of the 14 competing methods, some populated the top ranks of the different performance measures considerably more than others (Fig. 3). Counting the number of top 3 occurrences leads to the conclusion that methods 5, 1 and 2 (in this order) were most accurate overall. However, this approach naturally disfavors methods for which only partial results were submitted, and a closer look reveals that some of these methods actually performed better for specific conditions. Examples include method 3, which performed best in terms of α , β , JSC and JSC $_{\theta}$ for the higher-SNR data of scenario 3 and was among the top 3 best methods for many cases of scenario 1 (the only two scenarios for which results were submitted for this method); method 4, which performed best in terms of α and β for the higher-SNR data of scenario 2 (the only data for which results were submitted for this method) and in most cases was also the best in terms of RMSE for that data; method 7, which showed the best performance in terms of both JSC measures for the higher-SNR data of scenario 1; method 8, which, particularly in terms of α and β , performed best or second best for the

(Fig. 2 and Supplementary Table 2) can be used either to anticipate the success rate of automated particle tracking given the image quality or to determine the image quality required to assure a desired performance level according to the different criteria.

Analysis of biophysical measures

Although we used a comprehensive set of quantitative measures based on concepts also used in other fields, other, more specific measures might be desirable for specific biophysical analyses. Such measures can be easily applied retrospectively, as all results from our study are publicly available. To illustrate this, we performed additional analyses on the tracking results of the methods included in this study. Specifically, for each method and for each case for which results were submitted for that method, we computed the mean-squared displacement (MSD) for a representative range of time intervals (Supplementary Table 6). The resulting MSD curves (Supplementary Figs. 1–4) represent the estimated dynamic behavior of the particles. Generally, these results confirmed our finding that accuracy increases with increasing SNR and decreasing particle density. Furthermore, we observed that if particle motion is more purely diffusive (as in the vesicle scenarios of our study) rather than directed (as in the microtubule scenarios), most methods are less sensitive to SNR in estimating the MSD and yield good estimates also for SNR as low as 2 or even 1. This is to be expected, as in that case the displacements from one time point to the next are uncorrelated, and track switching errors have much less impact if all particles are subject to the same diffusion process. We also observed that in the case of a directed motion component (all considered scenarios except the vesicle scenarios), there is a general tendency by many methods to underestimate the MSD. This may be explained by the fact that longer particle jumps are more likely to be missed (the tracking methods may be too restrictive) and that track switching errors bias the results toward diffusive motion over longer time scales (if we assume track directions to be random and uncorrelated). We found that, by and large, the top-performing methods (Fig. 3) also performed best in terms of MSD estimation for the indicated cases, reconfirming the suitability of the measures we used for the competition. Similar observations followed from analyzing the results of instantaneous velocity estimation (Supplementary Table 7 and Supplementary Figs. 5–8). Finally, our retrospective analysis of the distribution of localization errors (Supplementary Table 8 and Supplementary Figs. 9–12) support and enhance our conclusions above regarding the top-performing methods in terms of RMSE.

DISCUSSION

We acknowledge that our study was one possible comparison of particle tracking methods, and future studies may extend ours in any of its three main aspects: methods, data or measures. Regarding the first, by design our study was limited to those methods developed by teams willing to participate in the competition at the time it was held. Fortunately, many traditional as well as more sophisticated tracking methods were included, and we believe our study was representative of the present state of the art. Regarding data, our study was limited to computer simulations, as this allowed for a controlled analysis (based on absolute ground truth) of tracking performance as a function of different factors. Although we believe that we considered the

most important factors (dynamics, density and signal), additional factors could be modeled, such as nonuniform background (more cell like), particle shape and size (varying within images and over time), frame rate (relative to particle velocity) and photobleaching (effectively allowing a time-dependent SNR). However, not only would a full analysis of all these factors be hampered by the ‘curse of dimensionality’—that is, an increased difficulty and resource requirement—but we can also expect tracking methods to perform only worse in such (more complex) data; even with most of our data, no method performed anywhere near perfectly. The ultimate challenge remains to obtain real experimental image data with as-accurate-as-possible ground truth. It has been suggested, for example, to use piezo stage-controlled particle motion⁶⁰.

Notwithstanding inevitable practical constraints, we believe that the present study is a major step toward more objective comparison of particle tracking methods, yielding important results and lessons for future development and experimentation. We identified important factors affecting particle tracking in practice and developed software for computer simulation of challenging image data to analyze tracking performance as a function of these factors. We also identified important measures to quantitatively score estimated tracks with respect to ground-truth reference tracks and developed software to automatically compute them. The software tools are publicly available as part of this article and can be used or further extended by any of those who are interested in benchmarking their particle tracking methods. We mobilized the field and stimulated groups worldwide to compare their methods in an open competition to improve transparency for potential users of the methods. Finally, we used the competition framework to compare current state-of-the-art particle tracking methods, and we performed additional analyses to illustrate the possibility to retrospectively study the impact on specific biophysical parameters beyond those considered in the competition itself.

In closing this article we summarize the main lessons learned for users and developers. Our results indicate that, at present, there exists no universally best method for particle tracking. Users should be aware that a method reported to work for certain experiments may not be the right choice for their application. As we pointed out, it is advisable to use synthetic image data mimicking the real data at hand, both to find the best parameter settings of a given method and to assess its potential performance. To this end, the tools developed as part of our study will prove useful for a wide range of biological scenarios, and the presented results already enable users to anticipate the performance of the tested methods for their applications. Users should be especially cautious when the SNR of their images is considerably lower than 4 (with our definition of SNR), although in the case of more diffusive (rather than directed) particle motion, most methods are able to yield accurate estimations of dynamics even for lower SNR. In selecting a method, users should also bear in mind that methods based on multiframe and/or multitrack optimization schemes in the linking stage, as well as well-tuned motion models, are likely to perform better than methods using simple per-frame and per-particle nearest-neighbor approaches. Thus, although more sophisticated methods may be more difficult to comprehend and control, they may be worth the time investment. For developers, the importance of parameter tuning and making the best possible use of prior knowledge about the data emphasizes the need for domain modeling in computational image analysis and suggests

the use of learning-based tracking methods. Because none of the tested methods performed perfectly on any of the data, and real biological data can be even more complex, the quest for better particle tracking methods remains. The results of the present study will serve as a useful baseline for testing the performance of future methods.

METHODS

Methods and any associated references are available in the [online version of the paper](#).

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AUTHOR CONTRIBUTIONS

E.M. and J.-C.O.-M. conceived of the study and organized the corresponding workshop at the 2012 IEEE International Symposium on Biomedical Imaging. F.d.C., I.S. and N.C. designed and implemented the software for image generation and comparison of tracking results. F.d.C. generated the image data and ground truth used in the training phase. M.M. generated the image data and ground truth used in the competition phase. M.M. and C.O.d.S. verified the files submitted by the participants and performed the scoring and assessed the computational speed of all methods. M.M., N.C. and E.M. analyzed the results. The methods were designed, implemented, run, and described by the participating teams: method 1: I.F.S., Y.G. and J.C.; method 2: C.C. and S.C.; method 3: N.C., F.d.C. and J.-C.O.-M.; method 4: M.W. and A.R.C.; method 5: W.J.G. and K.R.; method 6: Y.K.; method 7: L.L., J.D., H.S. and Y.X.; method 8: K.E.G.M., J.J. and H.M.B.; method 9: P.P.-G.; method 10: P.R., C.K. and F.W.; method 11: I.S. and E.M.; method 12: J.-Y.T. and S.L.S.; method 13: J.W., K.C. and G.P.v.W.; and method 14: H.-W.D. and Y.-S.T. The article was written by E.M. with input from all authors.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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- Saxton, M.J. & Jacobson, K. Single-particle tracking: applications to membrane dynamics. *Annu. Rev. Biophys. Biomol. Struct.* **26**, 373–399 (1997).
- Akhmanova, A. & Steinmetz, M.O. Tracking the ends: a dynamic protein network controls the fate of microtubule tips. *Nat. Rev. Mol. Cell Biol.* **9**, 309–322 (2008).
- Berginski, M.E., Vitriol, E.A., Hahn, K.M. & Gomez, S.M. High-resolution quantification of focal adhesion spatiotemporal dynamics in living cells. *PLoS ONE* **6**, e22025 (2011).
- Brandenburg, B. & Zhuang, X. Virus trafficking—learning from single-virus tracking. *Nat. Rev. Microbiol.* **5**, 197–208 (2007).
- Jandt, U. & Zeng, A.-P. Modeling of intracellular transport and compartmentation. *Adv. Biochem. Eng. Biotechnol.* **127**, 221–249 (2012).
- Sinha, B. *et al.* Dynamic organization of chromatin assembly and transcription factories in living cells. *Methods Cell Biol.* **98**, 57–78 (2010).
- Agarwal, S. *et al.* ATP-dependent and independent functions of Rad54 in genome maintenance. *J. Cell Biol.* **192**, 735–750 (2011).
- Stephens, D.J. & Allan, V.J. Light microscopy techniques for live cell imaging. *Science* **300**, 82–86 (2003).
- Ji, N., Shroff, H., Zhong, H. & Betzig, E. Advances in the speed and resolution of light microscopy. *Curr. Opin. Neurobiol.* **18**, 605–616 (2008).
- Shaner, N.C., Steinbach, P.A. & Tsien, R.Y. A guide to choosing fluorescent proteins. *Nat. Methods* **2**, 905–909 (2005).
- Giepmans, B.N.G., Adams, S.R., Ellisman, M.H. & Tsien, R.Y. The fluorescent toolbox for assessing protein location and function. *Science* **312**, 217–224 (2006).
- Saxton, M.J. Single-particle tracking: connecting the dots. *Nat. Methods* **5**, 671–672 (2008).
- Genovesio, A. *et al.* Multiple particle tracking in 3-D+t microscopy: method and application to the tracking of endocytosed quantum dots. *IEEE Trans. Image Process.* **15**, 1062–1070 (2006).
- Smal, I., Draegestein, K., Galjart, N., Niessen, W. & Meijering, E. Particle filtering for multiple object tracking in dynamic fluorescence microscopy images: application to microtubule growth analysis. *IEEE Trans. Med. Imaging* **27**, 789–804 (2008).
- Kalaidzidis, Y. Multiple objects tracking in fluorescence microscopy. *J. Math. Biol.* **58**, 57–80 (2009).
- Meijering, E., Dzyubachyk, O. & Smal, I. Methods for cell and particle tracking. *Methods Enzymol.* **504**, 183–200 (2012).
- Meijering, E., Smal, I. & Danuser, G. Tracking in molecular bioimaging. *IEEE Signal Process. Mag.* **23**, 46–53 (2006).
- Kalaidzidis, Y. Intracellular objects tracking. *Eur. J. Cell Biol.* **86**, 569–578 (2007).
- Dorn, J.F., Danuser, G. & Yang, G. Computational processing and analysis of dynamic fluorescence image data. *Methods Cell Biol.* **85**, 497–538 (2008).
- Meijering, E., Dzyubachyk, O., Smal, I. & van Cappellen, W.A. Tracking in cell and developmental biology. *Semin. Cell Dev. Biol.* **20**, 894–902 (2009).
- Jaqaman, K. & Danuser, G. Computational image analysis of cellular dynamics: a case study based on particle tracking. *Cold Spring Harb. Protoc.* **2009**, pdb.top65 (2009).
- Rohr, K. *et al.* Tracking and quantitative analysis of dynamic movements of cells and particles. *Cold Spring Harb. Protoc.* **2010**, pdb.top80 (2010).
- Blackman, S. & Popoli, R. *Design and Analysis of Modern Tracking Systems* (Artech House, Norwood, Massachusetts, USA, 1999).
- Sonka, M., Hlavac, V. & Boyle, R. *Image Processing, Analysis, and Machine Vision* 3rd edn. (Cengage Learning, Florence, Kentucky, USA, 2007).
- Cheezum, M.K., Walker, W.F. & Guilford, W.H. Quantitative comparison of algorithms for tracking single fluorescent particles. *Biophys. J.* **81**, 2378–2388 (2001).
- Carter, B.C., Shubeita, G.T. & Gross, S.P. Tracking single particles: a user-friendly quantitative evaluation. *Phys. Biol.* **2**, 60–72 (2005).
- Smal, I., Loog, M., Niessen, W. & Meijering, E. Quantitative comparison of spot detection methods in fluorescence microscopy. *IEEE Trans. Med. Imaging* **29**, 282–301 (2010).
- Ruusuvuori, P. *et al.* Evaluation of methods for detection of fluorescence labeled subcellular objects in microscope images. *BMC Bioinformatics* **11**, 248 (2010).
- Godinez, W.J. *et al.* Deterministic and probabilistic approaches for tracking virus particles in time-lapse fluorescence microscopy image sequences. *Med. Image Anal.* **13**, 325–342 (2009).
- Gillette, T.A., Brown, K.M., Svoboda, K., Liu, Y. & Ascoli, G.A. DIADeMchallenge.org: a compendium of resources fostering the continuous development of automated neuronal reconstruction. *Neuroinformatics* **9**, 303–304 (2011).
- Anonymous. Going for algorithm gold. *Nat. Methods* **5**, 659 (2008).
- Sbalzarini, I.F. & Koumoutsakos, P. Feature point tracking and trajectory analysis for video imaging in cell biology. *J. Struct. Biol.* **151**, 182–195 (2005).

33. Coraluppi, S. & Carthel, C. Recursive track fusion for multi-sensor surveillance. *Inf. Fusion* **5**, 23–33 (2004).
34. Coraluppi, S. & Carthel, C. Multi-stage multiple-hypothesis tracking. *J. Adv. Inf. Fusion* **6**, 57–67 (2011).
35. Olivo-Marin, J.-C. Extraction of spots in biological images using multiscale products. *Pattern Recognit.* **35**, 1989–1996 (2002).
36. Chenouard, N., Bloch, I. & Olivo-Marin, J.-C. Multiple hypothesis tracking in cluttered condition. in *Proc. Int. Conf. Image Proc.* 3621–3624 (IEEE, 2009).
37. Chenouard, N., Bloch, I. & Olivo-Marin, J.-C. Multiple hypothesis tracking in microscopy images. in *Proc. IEEE Int. Symp. Biomed. Imaging* 1346–1349 (IEEE, 2009).
38. Winter, M.R., Fang, C., Banker, G., Roysam, B. & Cohen, A.R. Axonal transport analysis using Multitemporal Association Tracking. *Int. J. Comput. Biol. Drug Des.* **5**, 35–48 (2012).
39. Winter, M. *et al.* Vertebrate neural stem cell segmentation, tracking and lineageing with validation and editing. *Nat. Protoc.* **6**, 1942–1952 (2011).
40. Godinez, W.J., Lampe, M., Eils, R., Müller, B. & Rohr, K. Tracking multiple particles in fluorescence microscopy images via probabilistic data association. in *Proc. IEEE Int. Symp. Biomed. Imaging* 1925–1928 (IEEE, 2011).
41. Rink, J., Ghigo, E., Kalaidzidis, Y. & Zerial, M. Rab conversion as a mechanism of progression from early to late endosomes. *Cell* **122**, 735–749 (2005).
42. Liang, L., Shen, H., De Camilli, P. & Duncan, J.S. Tracking clathrin coated pits with a multiple hypothesis based method. *Med. Image Comput. Assist. Interv.* **6362**, 315–322 (2010).
43. Yin, Z., Kanade, T. & Chen, M. Understanding the phase contrast optics to restore artifact-free microscopy images for segmentation. *Med. Image Anal.* **16**, 1047–1062 (2012).
44. Magnusson, K.E.G. & Jaldén, J. A batch algorithm using iterative application of the Viterbi algorithm to track cells and construct cell lineageing. in *Proc. IEEE Int. Symp. Biomed. Imaging* 382–385 (IEEE, 2012).
45. Husain, M., Boudier, T., Paul-Gilloteaux, P., Casuso, I. & Scheuring, S. Software for drift compensation, particle tracking and particle analysis of high-speed atomic force microscopy image series. *J. Mol. Recognit.* **25**, 292–298 (2012).
46. Casuso, I. *et al.* Characterization of the motion of membrane proteins using high-speed atomic force microscopy. *Nat. Nanotechnol.* **7**, 525–529 (2012).
47. Rao, A.R. & Schunck, B.G. Computing oriented texture fields. *CVGIP: Graph. Models Image Process.* **53**, 157–185 (1991).
48. Hager, G.D., Dewan, M. & Stewart, C.V. Multiple kernel tracking with SSD. in *Proc. IEEE Comput. Soc. Conf. Comput. Vis. Pattern Recognit.* 790–797 (IEEE, 2004).
49. Rousseeuw, P.J. & Leroy, A.M. *Robust Regression and Outlier Detection* (Wiley, Hoboken, New Jersey, USA, 2003).
50. Thompson, R.E., Larson, D.R. & Webb, W.W. Precise nanometer localization analysis for individual fluorescent probes. *Biophys. J.* **82**, 2775–2783 (2002).
51. Shafique, K. & Shah, M. A noniterative greedy algorithm for multiframe point correspondence. *IEEE Trans. Pattern Anal. Mach. Intell.* **27**, 51–65 (2005).
52. Jaqaman, K. *et al.* Robust single-particle tracking in live-cell time-lapse sequences. *Nat. Methods* **5**, 695–702 (2008).
53. Lowe, D.G. Distinctive image features from scale-invariant keypoints. *Int. J. Comput. Vis.* **60**, 91–110 (2004).
54. Crocker, J.C. & Grier, D.G. Methods of digital video microscopy for colloidal studies. *J. Colloid Interface Sci.* **179**, 298–310 (1996).
55. Celler, K., van Wezel, G.P. & Willemsse, J. Single particle tracking of dynamically localizing TatA complexes in *Streptomyces coelicolor*. *Biochem. Biophys. Res. Commun.* **438**, 38–42 (2013).
56. Ku, T.-C. *et al.* An automated tracking system to measure the dynamic properties of vesicles in living cells. *Microsc. Res. Tech.* **70**, 119–134 (2007).
57. Ku, T.-C., Kao, L.-S., Lin, C.-C. & Tsai, Y.-S. Morphological filter improve the efficiency of automated tracking of secretory vesicles with various dynamic properties. *Microsc. Res. Tech.* **72**, 639–649 (2009).
58. Huth, J. *et al.* Significantly improved precision of cell migration analysis in time-lapse video microscopy through use of a fully automated tracking system. *BMC Cell Biol.* **11**, 24 (2010).
59. Munkres, J. Algorithms for the assignment and transportation problems. *J. Soc. Ind. Appl. Math.* **5**, 32–38 (1957).
60. Saxton, M.J. Wanted: a positive control for anomalous subdiffusion. *Biophys. J.* **103**, 2411–2422 (2012).

ONLINE METHODS

Software implementations. The software for fully automated generation of the simulated image data used in this study and the software for computation of the performance measures (**Supplementary Note 4**) were written in the Java programming language as plug-ins for the open bioimage informatics platform Icy⁶¹ (**Supplementary Software**). Software implementations of the particle tracking methods of the participating teams (**Supplementary Note 1**) were written using various programming languages and platforms, including Java (stand-alone modules or plug-ins for ImageJ/Fiji⁶² or Icy), C++ (provided as source code or executable) and Matlab (MathWorks).

Analysis of performance results. For each tracking method and each performance measure, 48 values could in principle be computed, corresponding to the 48 data cases (different combinations of particle dynamics, densities and signal levels). However, not all teams submitted tracking results for all cases, which ruled out the possibility to perform an overall comparison and ranking of the different methods based on all cases. We observed that teams who did not apply their method to all 48 cases generally focused on one or more of the four dynamics scenarios representing different biological applications, but even per scenario not all teams applied their method to all pertaining cases. Therefore, we decided to rank the methods according to best performance per measure and per data case (**Fig. 3** and **Supplementary Table 2**).

Verification of tracking results. Minor differences between the originally submitted tracking results and the verified results

were to be expected because some of the software tools were converted to another platform to allow execution on the single evaluation system, and some methods were probabilistic in nature. Therefore, for each method, differences were considered acceptable (reproducible) if their means for each of α , β , JSC and JSC_{θ} were within 3% and the RMSE was within 0.5 pixel. In the vast majority of cases, the differences were acceptable, and the larger differences in some cases could be traced back to bug fixes and minor improvements in the software or parameter settings used for verification as compared to the original versions. In very few instances the results could not be verified owing to hardware or software limitations (**Supplementary Table 3**). For the analysis, the performance values computed from the originally submitted tracking results, not the verified results, were used.

Scoring of computation times. Computation times of all methods were measured on a single workstation (64-bit Intel Xeon X5550 2.67 GHz processor with 24 GB of RAM and running Microsoft Windows 7 Professional or Linux Fedora 16) to allow a fair comparison. We timed only those cases for which tracking results were submitted and verified. Similarly to the analysis of the accuracy performance measures, we ranked the methods according to best timing per data case (**Fig. 3** and **Supplementary Table 4**).

61. de Chaumont, F. *et al.* Icy: an open bioimage informatics platform for extended reproducible research. *Nat. Methods* **9**, 690–696 (2012).
62. Schindelin, J. *et al.* Fiji: an open-source platform for biological-image analysis. *Nat. Methods* **9**, 676–682 (2012).