The Nuclear Proteome of a Vertebrate

Graphical Abstract

Highlights

- Nucleocytoplasmic partitioning was quantified for 9,000 proteins in *Xenopus* oocytes
- Partitioned proteins have a native molecular weight larger than ~100 kDa
- Only a small fraction of proteins respond to Exportin 1 inhibition
- Passive retention is the dominant mechanism for the maintenance of the nuclear proteome

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In Brief

Wühr et al. quantify the nucleocytoplasmic partitioning for ~9,000 proteins in the *Xenopus* oocyte. Most proteins localize almost exclusively to nucleus or cytoplasm. Proteome-wide analysis of native protein size reveals that the distinct composition of nucleus and cytoplasm is primarily maintained by retention of proteins in large complexes.
The Nuclear Proteome of a Vertebrate

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SUMMARY

The composition of the nucleolus determines the behavior of key processes such as transcription, yet there is still no reliable and quantitative resource of nuclear proteins. Furthermore, it is still unclear how the distinct nuclear and cytoplasmic compositions are maintained. To describe the nuclear proteome quantitatively, we isolated the large nuclei of frog oocytes via microdissection and measured the nucleo-cytoplasmic partitioning of ~9,000 proteins by mass spectrometry. Most proteins localize entirely to either nucleus or cytoplasm; only ~17% partition equally. A protein’s native size in a complex, but not polypeptide molecular weight, is predictive of localization: partitioned proteins exhibit native sizes larger than ~100 kDa, whereas natively smaller proteins are equidistributed. To evaluate the role of nuclear export in maintaining localization, we inhibited Exportin 1. This resulted in the expected re-localization of proteins toward the nucleus, but only 3% of the proteome was affected. Thus, complex assembly and passive retention, rather than continuous active transport, is the dominant mechanism for the maintenance of nuclear and cytoplasmic proteomes.

INTRODUCTION

The organization of cells into membrane-enclosed compartments (i.e., organelles), each housing a characteristic set of macromolecules, is one of the foundations of complex, eukaryotic life [1]. Access of proteins to the nucleus is often highly regulated and controls critical steps in development, stress response, and general cell signaling [2].

Molecular traffic between nucleus and cytoplasm is routed through nuclear pore complexes (NPCs) embedded in the nuclear envelope [3]. These pores are permeable to ions, metabolites, and small proteins (reported to be up to ~40 kDa in molecular weight) but do not allow larger macromolecules to pass efficiently unless they are bound by nuclear transport receptors (also called karyopherins) that include importins and exportins [4–6]. Their activity is rendered directional and energy dependent by the coupling of transport to the RanGTPase system [7].

Despite the central role of the nucleus in multicellular biology, its protein content has never been satisfactorily cataloged, nor has the proteome’s nucleocytoplasmic partitioning been quantified systematically. This is at least partly due to the fact that efficient separation of nuclear and cytoplasmic material remains a serious challenge: the time required for cell fractionation is long compared to the time it takes some nuclear proteins to escape via diffusion [4, 8]. Furthermore, the relative quantification of protein abundance on a proteome-wide scale is only recently possible thanks to advances in mass spectrometry.

How the nuclear proteome is established during nuclear formation and subsequently maintained during interphase remains an open question. In animals and plants, the nucleus disassembles during mitosis and is rebuilt thereafter. Nuclear import plays a fundamental role in establishing nuclear composition [9, 10]. Throughout interphase, which can last many years in some somatic cells, nuclear composition has to be maintained. This is a challenge as proteins smaller than ~40 kDa in molecular weight can pass nuclear pores freely. Diffusion of larger proteins is restricted, but not completely prevented. Ultimately, this would lead to intermixing of nuclear and cytoplasmic contents. Continuous nuclear export has been shown to keep cytosolic proteins out of the nucleus [11]. As an alternative but not incompatible mechanism, proteins may bind large structures like DNA or assemble into large protein complexes, thereby practically preventing their diffusion through the pores. For example, antibody fragments directed against histones remain in the nucleus even though they lack a nuclear localization signal [12]. The contributions of active transport and passive retention to the maintenance of distinct nuclear and cytoplasmic proteomes have never been systematically investigated on the level of the proteome. While retention makes sense for proteins tightly bound to chromatin, it is not at all clear that the soluble contents of the nucleus (or the cytoplasm) can be maintained that way.

Our initial goal was to use a simple but reliable method of nuclear purification, the manual isolation of the large nuclei of the frog oocyte, to generate a reliable catalog of nuclear and cytosolic proteins. These could be accurately quantified using two recently developed methods of quantitative proteomics. Since the state of complex formation would be concentration dependent, we assessed the native molecular weight of proteins in undiluted cytosol and analyzed how nucleocytoplasmic protein localization is affected by inhibition of the cell’s major nuclear export pathway. This allowed us to address fundamental questions of how the nuclear content is maintained.

RESULTS

Proteome-wide Quantification of Nucleocytoplasmic Partitioning

Among organelles of eukaryotic cells, the nucleus is unique in not having a continuous membrane segregating its internal contents...
from the cytosol. In isolation procedures performed with tissue culture cells, soluble nuclear proteins could diffuse out through the nuclear pores, as well as through any other breaches in the membrane adventitiously generated by detergent or mechanical isolation. These problems may have contributed to poor agreement about just what is a nuclear protein. A remarkable exception to the problems of nuclear isolation is the microdissection of the millimeter-sized oocytes of amphibians. The giant nuclei (~400 μm diameter) of Xenopus laevis oocytes can be isolated manually, which minimizes loss of material due to comparatively quick isolation and the much longer time (about 10,000-fold) it would take proteins to diffuse on this length scale compared to somatic nuclei (Movie S1) [8]. To quantify nucleocytoplasmic protein partitioning in a proteome-wide manner, we determined relative nucleocytoplasmic protein concentrations in biological and technical triplicates using two different methods of accurate multiplexed proteomics (MultiNotch MS3 and TMTC) [13, 14] (Figures 1A and S1) along with our recently described genome-free proteomics approach [15]. To further control for protein leakage, we performed nuclear isolation for experiment 3 under mineral oil. We also demonstrated that the leakage of GST-tagged NLS-GFP out of the nucleus is much slower than nuclear isolation (Movies S1 and S2). For each quantified protein, we calculated the relative nuclear concentration (RNC), defined as the ratio of concentrations in the nucleus to the concentrations in nucleus plus the cytoplasm (Figures S1B and S1C). The RNC values obtained from the three replicates agree well, with an R² of at least 0.94 (Figures 1B and S2A). Altogether, we quantified the RNCs for 9,262 proteins (Figure S2B and Table S1A).

The RNC histogram revealed a distinct trimodal distribution: most proteins are localized almost exclusively to either the nucleus or cytoplasm, whereas a smaller third subset is nearly equally distributed (Figure 1C). When we used RNC values of 1/3 and 2/3 for discrimination, we quantified 55% of proteins as cytoplasmic, 17% as equidistributed, and 27% as nuclear. To compare and integrate our measurements with available metadata, which is typically human, we mapped the frog proteins to human homologs via a bidirectional best blast hit relationship (Figures 1B and S2). For each quantitative proteomics approach, we combined false discovery rate (FDR) <1%) that were observed in combined false discovery rate (FDR) <1%).
Correlation of Nucleocytoplasmic Partitioning and Native Molecular Weight

Our dataset allowed us to test the importance of the mechanisms proposed to be involved in nucleocytoplasmic partitioning. Two mechanisms have been suggested: first, some proteins may be retained in the nucleus or cytoplasm by virtue of their large hydrodynamic radii, which would impede movement through the nuclear pores [4, 21]; and second, continuous (energy-dependent) nuclear transport might be required to reverse the inevitable intermixing of nuclear and cytoplasmic proteins that would result in free diffusion through pores [11]. Of course, the cell employs both mechanisms to maintain nuclear and cytoplasmic composition, but their relative contribution has never been assessed. We were then in a position to evaluate these models directly at the proteome-wide level.

To test whether partitioned proteins are preferentially large, whereas equidistributed proteins tend to be small, we first compared the polypeptide molecular weight for cytoplasmic, equidistributed, and nuclear proteins. We found only a modest overrepresentation of low-molecular-weight proteins (<40 kDa) in the equidistributed fraction. In fact, many such proteins are either entirely nuclear or completely cytoplasmic (Figure 2A). Yet polypeptide mass is not a good indicator for the capacity to diffuse through nuclear pores. Rather, the native molecular weight of a protein, which considers whether a polypeptide chain might assemble into large complexes with other proteins or nucleic acids, is the much more appropriate measure. Although a number of distinct stoichiometric complexes are now known [22], our knowledge is likely to be far from comprehensive, and weaker and less specific assemblies, some of which would require the high concentration found in the cytosol, are generally elusive.

To determine whether the native size of proteins offered better discrimination between equidistributed proteins and those that are localized to either nucleus or cytoplasm, we developed a proteome-wide approach for estimating native protein size. We prepared undiluted frog egg extract by centrifugal crushing of packed eggs to minimize dilution of cellular contents, as such dilution might perturb complex formation. Unlike typical cell lysates, egg extract is still “alive” by many criteria: it can form metaphase spindles [23], cycle between interphase and mitosis [24], and form nuclei [25]. We then centrifuged the extract through protein filters of two molecular weight cutoffs (30 kDa and 100 kDa, respectively) and compared the input and filtered material by quantitative proteomics (Figure 2B). These filters do not give binary fractionation; rather, they admit proteins to an extent that varies continuously with molecular weight like many gel filtration materials. Thus, the degree of filtration yields graded information about the native size of a protein or complex.

To integrate the information from both filtration steps into a single value, we projected each data point onto a spline [26] and obtained a proxy for native size (Figure 2C). Comparison of this proxy against the known native molecular weight of proteins and protein complexes reported in the literature (Table S1B) revealed excellent correlation (R² of 0.95; Figure 2D). This allowed us to estimate the native molecular weight for ~3,500 proteins (Table S1C). This filtration-based approach should be generally applicable to investigate the formation of protein complexes and their dynamics in cell extracts.

Many proteins exhibited a much larger native molecular weight than predicted by their mere polypeptide molecular weight (Figure 2E). For example, small proteins in the anaphase-promoting complex, the proteasome, or the ribosome migrated with a molecular weight of more than 250 kDa, the upper size limit that we could resolve with the filters used (Table S1C). Although we saw only a weak correlation of polypeptide molecular weight and RNC (Figure 2A), the native molecular weight revealed a clear pattern of subcellular localization based on size (Figure 2F): essentially all natively small proteins are nearly equilibrated between nucleus and cytoplasm (RNC ~0.5). In contrast, most natively large proteins preferentially segregate either to the nucleus or cytoplasm, with some important exceptions (see below). The observed transition is gradual and occurs at approximately 100 kDa. This is larger than the reported size exclusion limit of NPCs (~40 kDa) [6]. We do not understand this discrepancy. It is possible that the functional size exclusion limit of NPCs is larger than the limit measured previously in short-term experiments [4]; over longer time periods, larger proteins may equilibrate. Alternatively, oocyte NPCs might be more permeable than those of somatic nuclei. Furthermore, although the literature typically reports an ~40 kDa cutoff, some studies have reported significantly larger cutoffs up to ~150 kDa [27, 28]. Nevertheless, our data strongly indicate that size exclusion by the NPC could maintain nucleocytoplasmic partitioning by preventing free diffusion of proteins and protein complexes larger than 100 kDa. That we observe hardly any small but partitioned proteins suggests that the cell does not typically spend transport receptor binding capacity and energy to maintain a nucleocytoplasmic concentration gradient for proteins that would diffuse rapidly through the nuclear pore.

Although most natively large proteins preferentially localize to one side of the nuclear membrane or the other, there is a small set of equipartitioned and natively large proteins, which can be seen in Figure 2F as the peak at RNC ~0.5 and native molecular weight >250 kDa. This set includes members of highly studied...
complexes like the anaphase-promoting complex (APC/C) and the proteasome (Figure 2G). We suspect that some undiscovered mechanism equipartitions these large complexes.

**Effect of Exportin 1 Inhibition on Nucleocytoplasmic Protein Partitioning**

It was proposed that continuous, energy-dependent nuclear export is required to keep cytoplasmic proteins out of the nucleus [11]. The nuclear export receptor Exportin 1 (CRM1) has been suggested to play a major role in the maintenance of nuclear identity and is believed to be the exportin with the most diverse cargo range [11, 29]. To assess the contribution of Exportin 1-mediated nuclear export to the maintenance of nuclear composition, we inhibited Exportin 1 with Leptomycin B (LMB) [30] and monitored nucleocytoplasmic protein distribution over time (Figure 3A). The vast majority of proteins quantified in
both replicates (6,411 out of 6,639; 97%) did not change their localization significantly, even after 24 hr of LMB treatment (Figures 3B and 3C and Table S1D). LMB was clearly effective as the remaining 3% of the proteins shift their RNC significantly toward the nucleus. Although in our experiment Exportin 1 does not seem to be required to keep the bulk of cytoplasmic proteins out of the nucleus, it is likely that its activity establishes these localization patterns initially, i.e., when nuclei re-assemble after mitosis. It is also likely that Exportin 1 is required to maintain cytosolic protein localization over very long timescales. The proteins that did re-localize after LMB application are interesting (Figure 3D). Most subunits of the equipartitioned APC/C moved toward the nucleus (Figure 3E), consistent with an active role of Exportin 1 in their equilibration, presumably in conjunction with an importin. In contrast, proteasome subunits did not respond to LMB (Figure 3E), indicating that different mechanisms operate here. Overall, we identified only 226 proteins that shifted localization significantly (1% FDR) toward the nucleus after inhibition of Exportin 1 (Figure 3C and Table S1D). Of these candidate Exportin 1 substrates, 187 have not previously been identified as Exportin 1 substrates (Figure S3A). We further characterized some of the LMB responders in human tissue culture cells (Table S2). Notably, we saw no sign of native size dependence in this response to LMB (Figure S3B).
that the vast majority of partitioned proteins are natively large mitochondrial proteins as nuclear (Figure S4)[41]. The finding allows these proteins to equilibrate between nucleus and cytoplasm. Nearly all concentrations in nucleus and cytoplasm. Diffusion through nuclear pores permits the passage of small molecules but restricts that of larger ones. We observed that the vast concentration of proteins exclusive to nucleus or cytoplasm can be estimated at ~2 mM [15]. This is ~200-fold higher than the estimated ~10 μM of all import and export receptors found in the oocyte, as calculated from the same source (Figure S3C). It seems to be an inescapable conclusion that a protein-autonomous mechanism such as passive retention is required to maintain nuclear composition in eukaryotic cells. We expect this to be true also for smaller somatic cells. However, it will be important to test this hypothesis experimentally.

Our conclusions do not diminish the importance of active nuclear transport in nucleocytoplasmic compartmentation: there is no doubt that import and export are required to segregate clear from cytoplasmic contents after mitosis, when the nucleus re-forms [9, 10, 36]. However, how much passive, merely size-dependent compartmentation mechanism contributes to the maintenance of pre-established localization was unknown. Is active nuclear transport at all required to separate nucleus and cytoplasm during interphase [11]? It surely will be vital to re-localize large complexes that can diffuse appreciably through NPCs over very long timescales, i.e., for cells with long interphases or post-mitosis [43]. Large complexes might also disassemble over time, allowing their smaller components access to their non-steady-state compartment. Furthermore, normally cytosolic proteins that fail to assemble into their native complexes after their biosynthesis might enter the nucleus by diffusion or active import. This is the case for poly-basic proteins (such as RNA-binding translation factors), whose charged domains often act as cryptic nuclear import signals [11]. In fact, importins operate as chaperones for exposed basic domains [44].

In this study, we did not analyze post-translational modifications like phosphorylation. For some proteins, we might have inadvertently averaged the subcellular localization of multiple distinct protein species. With quantitative phosphoproteomics [45], the role of phosphorylation on the proteome’s subcellular localization could be studied systematically.

Perhaps most surprisingly, our work revealed that the majority of the cell’s small proteins are found in complexes greater than 100 kDa in molecular weight. This seems to contradict biochemical experience. However, in such experiments, dilution and fractionation could easily dissociate large molecular assemblies. Furthermore, small proteins are easiest to purify while fractions found in large assemblies may be easily missed. Our results raise the question of whether protein-protein interactions at concentrations of ~100 mg/ml may enable many interactions that are simply not seen in in vitro conditions. There is anecdotal evidence in many cases where concentrated extracts diluted even 2- or 3-fold fail to carry out complex processes, like spindle formation, nuclear assembly, and cell-cycle progression. This aspect of the conclusion in this study, which after all assays protein distributions under native cellular conditions, warrants further study.

Finally, though we have stressed the generality of these mechanisms of nucleocytoplasmic partitioning, there will undoubtedly
be differences between oocytes and somatic cells. The nuclear proteins identified in this study appear to be mostly common to all cell types, but some are known to be special to the oocyte nucleus, for example, those involved in maintaining chromosomes for months in diplotene stage or those that enable the oocyte to reprogram somatic nuclei to totipotency [46]. While the comparison among different cell types could also be done via imaging methods, this would be very labor intensive and time consuming. Recently, very quick nuclear isolation methods for somatic cells have been developed [47]. Combining these with the quantitative proteomics analysis described here might be a promising strategy for nuclear proteome analysis of somatic cells.

**EXPERIMENTAL PROCEDURES**

**Nuclear Isolation**

The research with *X. laevis* was performed under the oversight of the Harvard Medical Area Institutional Animal Care and Use Committee. Isolation of *X. laevis* oocytes was done essentially as previously described [48]. J line (National Xenopus Resource Center) females were anesthetized with 0.2% Tricaine, and ovary lobes were surgically removed under sterile conditions. Oocytes were manually defolliculated and maintained in OCM (320 ml sterile water, 480 ml Lieblovitz medium [L-15] with glutamine [Sigma], 0.32 g BSA [Sigma], and 4 ml penicillin/streptomycin; the pH was adjusted to 7.7 with NaOH). Oocytes were allowed to recover overnight before the experiments. Before sample collection, oocytes were washed three times with MMR (0.1 M NaCl, 2 mM KCl, 1 mM MgSO4, 2 mM CaCl2, 5 mM HEPES [pH 7.8], and 0.1 mM EDTA) to remove BSA. For experiments LMB-1 and LMB-2, nuclei were isolated in MMR, and for experiment RNC-TMTC, nuclei were isolated under mineral oil (Sigma). For experiments LMB-1 and LMB-2, oocytes were transferred into MMR with 200 mM LMB. For each time experiment condition, 40–50 oocytes were separated into nucleus and cytoplasm and immediately frozen on dry ice. The control II for LMB-2 was collected, without drug treatment, and after 24 hr in LMB, samples were collected to control for effects solely due to time outside the oocyte. For confirmation of cell viability after 24 hr in LMB, their ability to respond to 3 nM progesterone was assayed (data not shown) [49]. Untreated cells were marked with Nile Blue and co-imaged [48]. Samples were lysed with 250 mM sucrose, 1% NP40 Substitute (Sigma), 5 mM EDTA (pH 7.2), 1 Roche Complete mini tablet (EDTA free), 20 mM HEPES (pH 7.2), 10 μM combetastatin A4, and 10 μM Cytochalasin D [15]. Lysate was vortexed at maximum speed for 10 s, pipetted ten times up and down with a 200 μl pipette tip, incubated on ice for 10 min, and again vortexed for 10 s. Lysates were clarified by centrifugation at 7,500 g at 4°C for 4 min in a tabletop centrifuge. After gentle flicking to resuspend lipids, supernatant was removed and used for further analysis. For the GFP-NLS leakage experiment (Movie S2), 10 μl of 28mg/ml of GST-tagged NLS-GFP (a kind gift of Daniel Levy) were injected into stage-VI oocytes. After ~24 hr, nuclei were isolated manually, and one picture was taken with bright-field illumination under a dissection microscope (for Movie S2, this picture was replicated and shown as t = 0.0 min). After the switch to fluorescent imaging, the leakage of GFP-NLS out of the nucleus was followed in 10 s intervals.

**Filter Percolation Experiment**

*Xenopus* egg extract was prepared as previously described [23]. Extract was released into interphase by addition of 0.4 mM Ca and incubated for 20 min at room temperature. Aliquots were flash frozen for further analysis. In technical duplicates, 200 μl of extract were added to Amicon Ultra-0.5 ml Centrifugal Filter Units with 30 kDa nominal molecular weight cutoff (Millipore), and 90 μl were added to Amicon Ultra-0.5 ml Centrifugal Filter Units with 100 kDa nominal molecular weight cutoff (Millipore). Filters were centrifuged for 30 min at 20°C at 5,000 g. The ~65 μl of 30 kDa percolate were frozen for further analysis. The ~32 μl of 100 kDa percolate were also frozen for further analysis. 0.8 μl of crude extract, 11 μl of 100 kDa filtrate, and 30 μl of 30kDa filtrate were used for mass spectrometry (MS) analysis.

**Data Analysis for Nucleocytoplasmic Partitioning Experiments**

Human gene symbols were assigned to all sequences based on reciprocal best BLAST hit against human proteins available from UniProt as previously described [19]. The ratio of nuclear to cytoplasmic content that match the gene symbols of equidistributed proteins (PFN1, ACTB, MDH1, TP1, PGK1, GRHPR, HBZ, ALOXE3, GSTO1, TALDO1, HSPA1A, FAM115C, GSTM1, FABP4, SOD1, and CFL1) were calculated for each experimental condition. For correction of pipetting errors, the nuclear signal from the corresponding condition was divided by this ratio. For LMB-2, the two controls were averaged to proteins were filtered out if the experiment 1 RNC result was more than 0.85 lower. Total for 24 hr LMB) disagreed by more than four average SDs, the protein was not quantified. Furthermore, proteins were filtered out if the RNC value of the 12 hr time point was more than four SDs outside the control or 24 hr time point. For experiment LMB-1, the protein was filtered out if the control or 24 hr time point. Importantly, for all filtering conditions, we did not make any assumptions about the directionality of the movement. For the final LMB responders, we only considered proteins which were quantified successfully in both LMB experiments.

**Data Analysis for LMB experiments**

The ratio of nuclear to cytoplasmic content that match gene symbols of equidistributed proteins (PFN1, ACTB, MDH1, TP1, PGK1, GRHPR, HBZ, ALOXE3, GSTO1, TALDO1, HSPA1A, FAM115C, GSTM1, FABP4, SOD1, and CFL1) was calculated for each experimental condition. For correction of pipetting errors, the nuclear signal from the corresponding condition was divided by this ratio as described above. Because slight errors in normalization would result in a large number of false-positive responders to LMB, we further normalized each condition with LMB and the second control in experiment 2, so that the median signal was equivalent to the corresponding control. Note that this will most likely lead to a slight underestimation for the actual move of proteins toward the nucleus. In the LMB-2 experiment, when RNC values between biological replicates (2× control, or 2× 24 hr LMB) disagreed by more than four average SDs, the protein was not quantified. Furthermore, proteins were filtered out if the RNC value of the 12 hr time point was more than four SDs outside the control or 24 hr time point. For experiment LMB-1, the protein was filtered out if the control or 24 hr time point. Importantly, for all filtering conditions, we did not make any assumptions about the directionality of the movement. For the final LMB responders, we only considered proteins which were quantified successfully in both LMB experiments.

**Data Analysis of Physiological Protein Size Measurement**

In technical duplicates, the ratios of flow-through over input were measured. The ratios were capped at 2 × 10−4 and 2 × 10−3 for the 30 kDa Filter and 2 × 10−3 and 2 × 10−2 for the 100 kDa filter, which is the approximate maximum dynamic range for these measurements. Protein ratios which differed by more than seven average SDs (in log-space) between biological replicates were filtered out. Rati o values between biological replicates were averaged in log-space. Theoretical protein size was estimated by multiplication of the number of amino acids with 110 Da. The splice was fit through data, using the spline function from Mathworks File Exchange, created by John D’Errico. We generated the spline out of third polynomial segments with four knots forced to be continuously increasing. To project the ratios for proteins onto the spine and measure its distance, we used the x2sn function from Mathworks Exchange, created by Juennjakob Dugge [26]. The result of “proxy for protein size” was plotted against native protein sizes from the literature [16, 50–52] (Table S1A). The correlation was used to estimate the physiological protein size in the experiment. We capped physiological protein sizes at minimally 2kDa and maximally 2kDa, which we estimate to be the maximum dynamic range for this experiment.

**ACCESSION NUMBERS**

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium [53] via the PRIDE partner repository with the dataset identifier PXD001297.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, two tables, and two movies and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2015.08.047.
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