A perspective of the dynamic structure of the nucleus explored at the single-molecule level

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Abstract Cellular life can be described as a dynamic equilibrium of a highly complex network of interacting molecules. For this reason, it is no longer sufficient to “only” know the identity of the participants in a cellular process, but questions such as where, when, and for how long also have to be addressed to understand the mechanism being investigated. Additionally, ensemble measurements may not sufficiently describe individual steps of molecular mobility, spatial-temporal resolution, kinetic parameters, and geographical mapping. It is vital to investigate where individual steps exactly occur to enhance our understanding of the living cell. The nucleus, home to many highly complex multi-order processes, such as replication, transcription, splicing, etc., provides a complicated, heterogeneous landscape. Its dynamics were studied to a new level of detail by fluorescence correlation spectroscopy (FCS). Single-molecule tracking, while still in its infancy in cell biology, is becoming a more and more attractive method to deduce key elements of this organelle. Here we discuss the potential of tracking single RNAs and proteins in the nucleus. Their dynamics, localization, and interaction rates will be vital to our understanding of cellular life. To demonstrate this, we provide a review of the HIV life cycle, which is an extremely elegant balance of nuclear and cytoplasmic functions and provides an opportunity to study mechanisms deeply integrated within the structure of the nucleus. In summary, we aim to present a specific, dynamic view of nuclear cellular life based on single molecule and FCS data and provide a prospective for the future.

Keywords live cell imaging · single-molecule tracking · nuclear structure · RNA mobility · protein mobility · HIV

Abbreviations
SMT Single-molecule tracking
FRAP Fluorescence recovery after photobleaching
FLIP Fluorescence loss in photobleaching
FCS Fluorescence correlation spectroscopy
STED Stimulated emission depletion
PALM Photo activation localization microscopy
3D-SIM 3D structured illumination microscopy
APD Avalanche photo diode
EMCCD Electron multiplied charged coupled device
poly(A) poly-adenylated RNA
RNA mRNP Messenger ribo-nucleo-particle
BR2 Balbiani Ring 2
U1 Uridine-rich small nuclear ribo-nucleo-snRNP particle
NLS Nuclear localization sequence
NES Nuclear export sequence
RNAPII RNA polymerase II
HIV Human immunodeficiency virus
RT Reverse transcriptase
IN Integrase
MA Matrix
NC Nucleocapsid
Vpr Viral protein R
PIC HIV preintegration complex
LTR Long terminal repeat
TAR Trans-activating responsive element in the HIV LTR
ARM Arginine-rich Motif
RRE Rev response element

Introduction

Our current picture of cell nuclear function is shifting from the idea of fully assembled molecular complexes towards molecules freely “roaming” in nuclear space (Pombo and Cook 1996; Phair and Misteli 2000; Bentley 2002). Here, transient interactions (protein–protein, protein–DNA, and protein–RNA) result in a stochastic but directional progression of multi-scale and multi-player processes (Elf et al. 2007; Zenklusen et al. 2008; Darzacq et al. 2009). While this change in paradigm challenges our view of the composition of multi-protein complexes, it also raises questions of how the nuclear space is organized, how the availability of factors is regulated and how interaction sites are found by the “right” interaction partners. While no membrane-bound compartments (such as in the cytoplasm) have been identified, the cell nucleus is functionally and spatially compartmentalized. Specific areas can be labeled, including the nucleolus, PcG bodies, nuclear speckles, Cajal bodies, Gems, cleavage bodies, perinucleolar compartments, the SAM68 nuclear body, PML bodies, etc. (Spector 2001) forming a “patterned” space. As a result, besides crowding effects also found in the cytoplasm, molecules in the nucleus encounter a highly complex, anisotropic landscape (Richter et al. 2007). To probe the dynamic functions of this landscape, an ensemble of technologies (such as FRAP, FLIP, confocal imaging etc.) have been applied to elucidate the dynamics of protein factors and RNA particles (Becker et al. 2002; Braga et al. 2007). Many functional protein factors have been found to vary in their immobile versus mobile states, as well as in their average off rates from potential binding sites (Misteli and Spector 1999; Grunwalde et al. 2006b; Gorski et al. 2008; Grunwald et al. 2008a). Similarly, a wide range of mobility distributions exist in RNA studies (Politz et al. 1998; Shaw-Tal et al. 2004; Siebrasse et al. 2008). Even nuclear bodies, that appear stationary in imaging, are actually dynamic complexes undergoing a steady exchange of their building blocks (Politz et al. 2006).

Taken together, these data draw a picture of a highly mobile environment with changing fractions of immobilized populations and a continual change in composition. Experimental access to probe this environment and its embedded processes at the molecular scale is limited as: nuclear organization is actively maintained and so can only be studied in living cells; ensemble technologies, being diffraction limited, fail when challenged by molecular interactions on the nanometer length scale; and the high dynamics of these often transient interactions demand collection of data in the milliseconds time scale, limiting the achievable signal-to-noise ratios.

Observing single molecules in the nucleus, a possibility?

While multiple strategies are employed today to challenge the resolution criteria (Hell 2003; Betzig et al. 2006; Hess et al. 2006; Rust et al. 2006), imaging of single-molecule signals can provide the position of the observed molecule with an accuracy that can overcome the resolution limit by a factor of ten in the living cell, and is in principle capable of providing nanometer precision (Bobroff 1986; Schmidt et al. 1996; Thompson et al. 2002; Yildiz et al. 1999). Resolution and localization precision are not interchangeable terms. Practically, nonlinearities (on/off states of fluorophores) are used to construct images with high spatial precision in techniques like Photo Activation Localization Microscopy (PALM) while quenching of emission states by stimulated emission (used in Stimulated Emission Depletion, STED) directly increases the resolution of the recorded signal. In PALM, the same routines are used to determine the position of a molecule as in single molecule imaging; however, the latter relies on the isolated observation of one molecule over time to reveal its dynamic property and can easily be adopted to live cell microscopy.
al. 2003). Continuous image acquisition also enables following a single protein or RNA over an extended amount of time utilizing the high localization accuracy for tracking rather than construction of a high resolution image. An immediate advantage is that no synchronization of cellular processes is needed as single-molecule tracking (SMT) is fast enough to observe the sequence of events in real time and synchronization is achieved during data analysis. Finally, by providing superb time and spatial information simultaneously, SMT is perhaps the tool of choice when it comes to investigating the dynamic landscape of molecular interactions in the living cell.

The question however is how fast SMT can be, or in other words, what is the fastest process observable by this technology? SMT has been shown to be fast enough to even follow freely moving molecules in an aqueous solution (Grunwald et al. 2006a). Measurements of nucleocytoplasmic transport have shown definitively that transient interactions in the living cell can be resolved on the nanometer length- and milliseconds time scales using SMT. In the same way, single protein factors and RNAs in the nucleus have been measured using inert test probes and functional protein factors (Goulian and Simon 2000; Kubitscheck et al. 2005; Grunwald et al. 2006b; Grunwald et al. 2008a). Compared with fluorescence correlation spectroscopy (FCS) or fluorescence recovery after photobleaching (FRAP), SMT provides the means to analyze multiple forms of mobility in heterogeneous environments. Even more, SMT enables observation of the molecular behavior over the whole field of view and not only within either the confocal volume of an FCS spot or the bleached area in FRAP. This makes the technique very valuable for studying intranuclear processes of proteins and RNAs in relation to defined nuclear structures.

However, observation of single molecules in living cells needs to overcome the problem of having too many overlapping signals from the labeled population of molecules. Adjustment of the number of labeled molecules is usually done by: providing small amounts of recombinant fluorescently labeled proteins (e.g., by microinjection), by careful titration of labeled ligand concentrations, by adjusting expression levels of fluorescent reporter molecules (e.g., using leaky promoters), or by signal amplification (e.g., DNA probes, molecular beacons or the MS2 system) (Bertrand et al. 1998; Bratu et al. 2003; Siebrasse et al. 2008). The resulting experimental design is then driven by optimizing the signal-to-noise ratio during data acquisition, usually by a combination of background reduction, signal multiplication on the detector (e.g., APDs or EMCCDs) and very sensitive detection at low read noise conditions. Finally, the observation time needs to be maximized, taking photobleaching rates, desirable signal intensity and applicable amounts of light into consideration.

While many pioneering studies used excitation energies in the range of several kilo Watts per square centimeter, cells and especially nuclear structures and processes are sensitive to high-photon fluxes (Dobrucki et al. 2007; Hoebe et al. 2007; Grunwald et al. 2008b; Heinze et al. 2009). In this context, calibrating the amount of light delivered to the cell is of crucial importance and can be done for broadband light sources as well as for laser based imaging (Grunwald et al. 2008b). Depending on the mobility of the observed molecules and the acquisition speed of the microscope system used, the acquired data is then analyzed based on the tracking of individual signals (see (Grunwald et al. 2008c) for a detailed review). Figure 1 details simple modifications that can be applied to commercially available equipment to improve performance for the detection of single molecules in living cells at appropriate time scales. In short, a laser illumination module (here shown in a customized home-built version, but also commercially available) is combined with a fast switching mechanism and a fast intensity regulator for the laser light and fiber coupled to the microscope. A stable stage with high precision and small drift is used to position samples above the objective lens and to allow for cell manipulation (e.g., microinjection) if needed. Lastly, a sensitive, fast camera, typically an EMCCD, is used for recording the single-molecule signal and is mounted as directly as possible onto the microscope to increase photon transmission. Either sequential imaging or a second detector is used to monitor a cellular reference structure.

To summarize, SMT and FCS provide the means to study single molecules or very small, locally resolved, ensembles of fluorescently labeled molecules in the living cell. In the following, we discuss our current view of the nuclear landscape, dynamical aspects of the nucleus and point out controversial areas that remain. The road to live cell imaging was laid down by confocal microscopy and today’s FCS implemen-
tions are building upon this technology (Eigen and Rigler 1994).

Nuclear mobility as seen by fluorescence correlation spectroscopy

The first experiment to provide a mobility profile of the nuclear space used FCS and fluorescein-labeled oligodeoxynucleotides (oligos) to measure intranuclear diffusion coefficients in living cells (Politz et al. 1998). Labeled poly-thymidine oligos, that are passively taken up by living cells, formed hybrids with the poly-adenylated tails of RNAs (poly(A)RNAs) in the nucleus, and two major fractions of diffusion rates were observed. Besides the free oligonucleotide diffusion, a slow moving fraction, interpreted as the labeled diffusive RNAs ($D_{\sim 10 \mu m^2/s}$), was detected. A third minor fraction was also detected with a 10-fold slower mobility rate (see Table 1). In another experiment, the effect of nuclear compartments on the movement of poly(A)RNAs in the nucleoplasm was compared to poly(A)RNA mobility in speckles (Politz et al. 2006). These values are similar to the mobility of the slower class reported in the first study (Table 1). Interestingly, temperature reduction from 37°C to 22°C had no

Fig. 1 An SMT microscope setup. A conventional wide-field epifluorescence microscope is custom-modified to optimize single molecule detection in biological samples. Illumination is laser based. Single-mode, low-noise lasers are the preferred choice. The excitation (laser) light is merged using dichromatic beam splitters and mirrors and coupled into an optical monomode fiber to deliver the light to the microscope. An acousto-optical tunable filter is used to attenuate laser power and switch illumination on and off with microsecond resolution. The fiber output can either be delivered as a collimated beam to the excitation tube lens or be imaged via a conjugated image plane into the objective’s back focal plane. Emitted fluorescence light, collected by a high numerical aperture objective, is separated from the excitation light by an appropriate dichromatic beam splitter, which reflects the three excitation lines into the sample, while the three fluorescence bands are transmitted to the detection system. Notch filters have shown excellent performance in excluding remaining excitation light. The microscope is mounted on a steel scaffold fixed to the optical table to allow easy access to the base port, where an EMCCD for single molecule tracking is mounted directly onto the microscope stand collecting signal in the primary image plane. A second camera, located at a side port, is used to detect reference images. A microinjection system allows delivery of fluorescently labeled probes into a living cell.
detectable effect on poly(A)RNA mobility, and was interpreted as meaning that movement between the nucleoplasm and speckles does not require metabolic energy, which is in stark contrast to a previous report (Calapez et al. 2002).

Probing the nuclear space with inert probe molecules, initially described by Grunwald et al. 2008a using SMT, was then used to explore the effects of molecular crowding on diffusion and binding of nuclear proteins in heterochromatin using FCS technology (Bancaud et al. 2009). Volume exclusion, diffusive barriers in more dense nuclear compartments and transient trapping in heterochromatin, as observed by Grunwald et al. 2008a, were confirmed. Using three independent chromatin interacting proteins as active probes, photoactivation experiments were carried out, and in euchromatin, diffusion and binding parameters were well explained by a diffusion limited model. However, when the same probes were tested in heterochromatin, where slower kinetics were expected (dependent on the heterochromatin to euchromatin abundance ratio), an unexpected biphasic mobility was observed that could not be explained by a diffusion reaction model or a random crowding model (Bancaud et al. 2009). Based on these results the authors deduced fractal geometry of chromatin as previously suggested (Wachsmuth et al. 2000).

### Single RNA tracking in the nucleus

Circumventing the small active measurement area and time averaging inherent to FCS, improvements in camera technology, and the development of RNA labeling systems, made imaging of messenger ribonucleoprotein particles (mRNPs) in living cells a possibility. One such labeling system is the MS2 system that consists of two components, i.e., a sequence of MS2 RNA stem-loops introduced into the RNA sequence of interest and a coat protein to which a fluorescent protein is fused. The coat protein binds to the RNA sequence with nanomolar affinity thereby generating a multiplexed signal sufficient for single molecule tracking (Stockley et al. 1995; Table 1)

| Reference | Construct | Method | Diffusion coefficient | Subcompartment |
|-----------|-----------|--------|-----------------------|----------------|
| (Grunwald et al. 2006b) | U1 snRNP bioactive | SMT | $D=0.5$ to $8 \ \mu m^2/s$ | Nucleoplasm, speckles |
| (Grunwald et al. 2008a) | Streptavidin-NLS inert | SMT | $D=0.8$ to $5 \ \mu m^2/s$ | Nucleoplasm, nucleolus, heterochromatin |
| (Speil and Kubitscheck 2010) | Ovalbumin inert | SMT | $D=0.5$ to $12 \ \mu m^2/s$ | Nucleoplasm, nucleolus |
| (Bancaud et al. 2009) | GFP-monomer | FCS | $D_1=29 \ \mu m^2/s$ | Euchromatin |
| | | | $D_1=87 \ \mu m^2/s$ | Aqueous solution |
| | | | $D_2=17 \ \mu m^2/s$ | Euchromatin |
| | | | $D_2=55 \ \mu m^2/s$ | Aqueous solution |
| | | | $D_3=7.7 \ \mu m^2/s$ | Nucleoplasm |
| (Politz et al. 1998) | Oligo(dT):poly(A) RNA | FCS | $D=1$ to $10 \ \mu m^2/s$ | Nucleoplasm |
| (Politz et al. 2006) | Oligo(dT):poly(A) RNA | FCS | $D_1=0.65 \ \mu m^2/s$ | Speckles |
| | | | $D_2=0.7 \ \mu m^2/s$ | Nucleoplasm |
| (Shav-Tal et al. 2004) | YFP-MS2 mRNP | FRAP | $D=0.09 \ \mu m^2/s$ | Nucleoplasm |
| (van den Bogaard and Tyagi 2009) | GFP-mRNA-96-mer | SMT | $D=0.04 \ \mu m^2/s$ | Nucleoplasm |
| | | | $D=0.033 \ \mu m^2/s$ | Nucleus |
| (Siebrasse et al. 2008) | DNA-labeled BR mRNP | SMT | $D=0.24$ to $4.0 \ \mu m^2/s$ | Salivary-gland cell nucleus |
| (Mor et al. 2010) | Different sized Dys mRNA | SMT | $D=0.005 \ \mu m^2/s$ | Nucleus |
| (Maertens et al. 2005) | HIV-IN | FCS | $D=10.5 \ \mu m^2/s$ | Nucleus |

*In these studies, diffusion coefficients were fixed to compare the relative amounts of the different mobility classes in different compartments. For details, see original literature*
Bertrand et al. 1998; Lago et al. 2001). To detect mRNPs in real time in living mammalian cells, 24 MS2 stem-loops were inserted downstream of a target RNA (Shav-Tal et al. 2004). Movements of individual mRNPs were followed for 100 frames using exposure times of 250 and 333 ms, respectively. The tracking criteria for mRNPs was set to more than eight frames of continuous observation, and the mean diffusion coefficient, in stark contrast to the FCS data cited above, was found to be 0.04 μm²/s at 37°C. This diffusion was also temperature dependent, as at room temperature the diffusion coefficient was reduced indicating that mRNPs released after transcription moved probabilistically through the nucleoplasm following simple laws of diffusion (Shav-Tal et al. 2004). When very large mRNPs were tracked in the nucleus (Dys-mRNP, ranging from 4.8 to 14 kb), an even lower mobility of mRNPs was found (D = ~0.005 μm²/s) (Mor et al. 2010). Interestingly, in this report, mRNPs were found to travel in chromatin free zones in a “channeled diffusion” manner with the accumulation of mobile particles in distinct chromatin free zones (Mor et al. 2010). While there is evidence of open spaces in the lamin network as revealed by three-dimensional structured illumination microscopy (3D-SIM) (Schermelleh et al. 2008), it is unclear whether these spaces are indeed channels and if they exist over the entire nucleus as predicted (Mor et al. 2010).

Another labeling approach uses molecular beacons for fluorescent labeling of mRNP (van den Bogaard and Tyagi 2009). A molecular beacon is composed of a stem hybrid that keeps the fluorophore close to a quencher rendering the probe initially non-fluorescent. When the probe hybridizes to a target sequence, the fluorophore separates from the quencher due to a conformational change, and becomes detectable. Sequence specific multiplexing of probes on a target RNA is possible, reaching up to ~100 probe copies/RNA, resulting in signal intensities of single mRNPs that can be detected as diffraction-limited spots in living cells. In an experiment where imaging was performed at 300 ms integration time, an average diffusion coefficient of 0.033 μm²/s was found for one half of the observed mRNPs, while the other half displayed a stationary behavior with a calculated diffusion coefficient of 0.0006 μm²/s. When the single mRNPs positions were mapped to a chromatin density image the stationary particles were located in high-density chromatin regions. In contrast, the mobile particles were preferably found in low chromatin density areas. ATP depletion increased the number of stalled particles but no strong effect on diffusion was found, again suggesting that mRNP movement is an ATP-independent process (van den Bogaard and Tyagi 2009).

All of these results are ~10–100-fold lower in their diffusion constants (see Table 1) compared to the FCS based mobility profile (Politz et al. 1998; Politz et al. 2006). In contrast to the above studies, however, when an insect model system (instead of a mammalian cell system) was used, i.e., balbiani ring (BR2) mRNAs (extremely large mRNAs found in the salivary glands of the Chironomus midge), four different mobile fractions, widely varying, were found approaching diffusion coefficients comparable to those found by FCS (Table 1) (Siebrasse et al. 2008). Interestingly, the distribution of diffusion coefficients is better explained as a probability distribution of mobile states that an mRNA molecule can adopt, rather than by the existence of distinct classes of differently mobile mRNAs, a first indication of which was already seen in protein mobility (see Fig. 2 and (Grunwald et al. 2006). In contrast to the above studies, however, when an insect model system (instead of a mammalian cell system) was used, i.e., balbiani ring (BR2) mRNAs (extremely large mRNAs found in the salivary glands of the Chironomus midge), four different mobile fractions, widely varying, were found approaching diffusion coefficients comparable to those found by FCS (Table 1) (Siebrasse et al. 2008). Interestingly, the distribution of diffusion coefficients is better explained as a probability distribution of mobile states that an mRNA molecule can adopt, rather than by the existence of distinct classes of differently mobile mRNAs, a first indication of which was already seen in protein mobility (see Fig. 2 and (Grunwald et al. 2006). In contrast to the above studies, however, when an insect model system (instead of a mammalian cell system) was used, i.e., balbiani ring (BR2) mRNAs (extremely large mRNAs found in the salivary glands of the Chironomus midge), four different mobile fractions, widely varying, were found approaching diffusion coefficients comparable to those found by FCS (Table 1) (Siebrasse et al. 2008). Interestingly, the distribution of diffusion coefficients is better explained as a probability distribution of mobile states that an mRNA molecule can adopt, rather than by the existence of distinct classes of differently mobile mRNAs, a first indication of which was already seen in protein mobility (see Fig. 2 and (Grunwald et al. 2006).
Single-molecule tracking in the nucleus

A

B

C

D

E

F

\[ D = 54 \pm 3 \mu m^2/s \]
Fluorescent labeling here was done either by microinjection of oligos complementary to BR2 mRNA or with a labeled-hrp36 protein that specifically binds BR2 mRNA (Siebrasse et al. 2008).

As there are such dramatic differences between the FCS data and tracking data of one group versus another, it is very clear that controversy exists in this field, which might be explained by clear technical differences in how the experiments were performed.

**Single protein tracking in the nucleus**

As we have seen above that RNA tracking is possible, it would be interesting to also track single proteins as proteins are the facilitators of many major cellular processes. One such study looked at the mobility of fluorescently labeled uridine-rich small nuclear ribonucleoproteins (U1 snRNPs), biologically active splicing factors (Grunwald et al. 2006b). GFP-labeled ASF/SF2 was used to mark nuclear speckles allowing direct comparison of U1 snRNP dynamics inside and outside of the nuclear speckles. Using high speed fluorescence microscopy, with frame rates of up to 200 Hz, no significant mobility was found for ∼80% of U1 snRNPs, possibly caused by molecular trapping in nuclear structures as well as immobilization in spliceosomes and post-splicing processes. A continuous range of mobility for U1 snRNPs, ranging from 0.5 to 8 μm²/s was found. The diffusion coefficient of 0.5 μm²/s corresponds to impeded uncomplexed single U1 snRNPs or higher organized spliceosome-complexes. Correspondingly, using FRAP experiments, a three to five fold reduction of the diffusion coefficient of larger molecules in the nuclei was also found (Gorski et al. 2006). From here we conclude that there is not just one kinetic condition for association and dissociation events of biologically active proteins in the nucleus.

How does the large immobile fraction of U1 snRNPs compare, however, to inert proteins? Tagged streptavidin coupled to a nuclear localization sequence (NLS) with a size of about 60 kDa, and a second probe, ovalbumin with a size of 45 kDa, were tracked in living cell nuclei (Grunwald et al. 2008a; Speil and Kubitscheck 2010). While streptavidin is not translocated by nuclear pores, primarily due to its size, ovalbumin likely passively transports into the nucleus. Using high speed fluorescence imaging at frame rates of up to 200 Hz, the streptavidin experiment succeeded in capturing even single traces of probe molecules and deduced a diffusion rate comparable to that of the inert GFP protein as seen by FCS (see Table 1 and Fig. 2) (Grunwald et al. 2008a; Bancaud et al. 2009). Different nuclear compartments affect the movement of inert proteins differently, but even in the nucleoplasm, two kinetic components (mobile and trapped) were observed, and the mobile fraction is widely spread over a wide range of diffusion coefficients. (The data could not be fitted satisfactorily assuming only one or two diffusing components, Table 1). Compared with the nucleoplasm (defined as space neither labeled by MeCP2 or ASF1 exclusion), proteins seemed to become trapped predominantly in pericentric heterochromatin resulting in fewer proteins moving freely (see Fig. 2). This trapping is correlated to the high abundance of chromatin fibers in this area. Even more exciting, and in contrast with the FCS study discussed above (Bancaud et al. 2009), proteins were trapped less frequently in the nucleolus compared to the nucleoplasm. While confocal imaging implies that the nucleolus excludes the test protein, the mobility data suggest minimal trapping in this compartment and combined with no retention of proteins at the compartment interface, exclusion turns out to be a dynamic effect, where mobile proteins leave the nucleolus frequently and fast (Grunwald et al. 2008a; Speil and Kubitscheck 2010). When ovalbumin was used, an inert protein that does not contain an NLS, comparable observations were made demonstrating the limited, if any, impact of the SV40 NLS-related electric charge on protein mobility in the nucleus.

**Imaging HIV in the living cell—a perspective**

An important question that can be addressed using single molecule tracking is the mobility of HIV (Human Immunodeficiency Virus) related factors, their interaction times and regulation of cellular distribution. We first provide a brief summary of the HIV lifecycle, including import and export processes of HIV genetic material through the nuclear membrane, which is a very elegant symphony between cytoplasmic and nuclear viral and host cell factors. Single molecule tracking
techniques might enable deeper understanding of the spatial-temporal dynamics of these processes thereby not only providing key information on very basic biological processes, but we surmise, also exposing vulnerabilities that could be targeted in the fight against this devastating disease.

The life cycle of HIV

The HIV virus is an RNA lentivirus with a genome comprising two single-RNA strands containing the full genetic information needed for viral replication in host cells (Luciw et al. 1996; Fields et al. 2007; Levy 2007). Through the process of fusion and endocytosis the virus enters the cell (Fig. 3), by binding its trimeric surface gp120 to a CD4 molecule present on the host cell (reviewed in (Gallo et al. 2003)). This binding induces a conformational change to a specific chemokine receptor (predominantly CCR5 or CXCR4 (Berger et al. 1999)), and this in turn induces a conformational change in gp41, another viral surface factor. Gp41 in its altered conformation opens up, thereby “spearing” the lipid bilayer of the target cell. This fusion of the viral and infected cell’s membranes results in the release of the HIV viral core into the cytoplasm (Chan et al. 1997; Tan et al. 1997; Weissenhorn et al. 1997; Caffrey et al. 1998). Viral uncoating then takes place (Briones et al. 2010) and the viral reverse transcription complex, comprising viral and host proteins, is assembled and cDNA synthesis begins. The result of reverse transcription is an HIV preintegration complex (PIC), which is then challenged with importing itself into the nucleus. (Lee et al. 2010). It is still unclear how the PIC imports into the nucleus (Dvorin et al. 2002; Bukrinsky 2004; Yamashita and Emerman 2005; Yamashita and Emerman 2006).

Following transport across the nuclear pore complex viral and host proteins again participate in integration of the viral genome into the host cell genome (Greene and Peterlin 2002). While this is primarily mediated by integrase (IN), which binds to the ends of the viral DNA, host proteins are also involved and required, though their precise functions remain unknown (Kalpana et al. 1994; Lee and Craigie 1994; Farnet and Bushman 1997; Li et al. 2000; Suzuki and Craigie 2002; Beitzel and Bushman 2003; Lin and Engelman 2003). LEDGF/p75 also associates with IN and has also been implicated in participating in its nuclear import and/or the integration process (Cherepanov et al. 2003; Maertens et al. 2003). As the HIV provirus can integrate into many different chromosomal locations in the cell it is tempting to explain viral latency (vs. transcriptional activity) as integration into repressed heterochromatin. While this is at least in part true, it is not the whole story (for reviews on HIV viral latency, see (Han et al. 2007; Coiras et al. 2009; Graci et al. 2009)).

In the case of a transcriptionally active integrated provirus, the 5’ LTR (long terminal repeat containing promoter elements (Taub e et al. 1999)) positions RNA polymerase II (RNAP II) at the site of initiation of transcription and is responsible for the assembly of the pre-initiation complex. While transcription can begin with these minimal components, RNAP II invariably fails to elongate efficiently (Kao et al. 1987). To achieve efficient elongation the viral
protein Tat is required which associates with host cyclin T1, which in turn recruits host Cdk9. Tat binds the 5′ bulge region of TAR (a 59-nucleotide stem-loop RNA element in the LTR) via its arginine-rich motif and recruitment of P-TEF-b (the complex formed by cyclin T and Cdk9) results in hyper-phosphorylation of the C-terminal domain of RNA polymerase II, thereby stimulating efficient transcriptional elongation (Zhou et al. 2003).

The integrated provirus, while only ∼9 kb long, then successfully expresses 15 distinct viral proteins, facilitated by an elegant and complex splicing mechanism that involves both complete and incomplete splicing (Frankel and Young 1998). When the mRNA is completely spliced (encoding for Nef, Tat, and Rev) it is rapidly transported into the cytoplasm and transcribed (Cullen 1998). When the mRNA, however, is singly spliced or unspliced, viral transcripts remain in the nucleus and are relatively stable (Luo et al. 1994; Powell et al. 1997). While the export of unspliced (or partially spliced) RNA in eukaryotic cells is usually prohibited, the virus overcomes this block by utilizing the viral protein REV (Custodio et al. 1999). The REV protein binds to an Rev response element (RRE) encoded in the viral RNA sequence and together with recruited host proteins manages to export the unspliced or partially spliced RNA in what is referred to as the REV-RRE-CRM1 export mechanism (reviewed in (Hope 1999)). While the partially (or singly) spliced viral transcripts encode the structural enzymatic accessory proteins, the unspliced RNA species constitute the genome of newly formed progeny virions, and export of these RNAs depend heavily on REV’s leucine-rich NES (nuclear-export sequence) as well as on host proteins, predominantly Ran GTPase (Fornerod et al. 1997).

As the viral components needed for new virion formation are assembled in the cytoplasm, shuttling to the plasma membrane (where budding will occur) takes place (reviewed in (Kaplan 2002; Li and Wild 2005; Mazze and Degreve 2006)). This process is highly reliant on the viral Gag polyprotein, but involves, once again, a number of host cell factors, without which the process does not occur (Dussupt et al. 2009). The terminal step in the budding reaction involves a second membrane fusion event for the virus, and this too is an orchestrated effort between the viral Gag polyprotein and host proteins (Kaplan 2002; Li and Wild 2005; Mazze and Degreve 2006).

The virion is then released and free to continue the vicious cycle of infection.

Where to from here? As becomes evident from the wealth of information already available, the past several decades of research have defined a large number of host cell proteins that influence every step of the viral life cycle (Ptak et al. 2008; Fu et al. 2009; Pinney et al. 2009). Initial imaging results on the localization of, for example CD4 and CCR5 on the cellular membrane (Steffens and Hope 2003; Steffens and Hope 2004), RNA distribution in virions (Chen et al. 2009), and the biogenesis of HIV virions at the plasma membrane (Jouvenet et al. 2008) have demonstrated the applicability of microscopy to the HIV field. However, a report that investigated IN mobility within the nucleus by FCS, also demonstrates the current limitations at the nanometer length- and milliseconds time-scale (Maertens et al. 2005). Clear examples of how SMT could be applied to the study of HIV include analysis of the import and export of viral RNA and proteins, e.g. the PIC complex, the interaction durations and sites for the REV-RRE-CRM1 complex, the distribution of genome integration sites or the interplay of TAT and RNAPII. While the above are good examples of what can be achieved with single-color SMT, investigating the interaction times and complex formation of, for instance HIV-RNA and Rev or IN protein, will require simultaneous multi-color SMT. The problem with this approach is that while in each individual color single molecules will be localized with high precision, the colocalization of these single-molecule positions will remain diffraction limited, hence in the range of more than 200 nm. While single-molecule motors, imaged directly on the cover glass in solution have been double labeled and registered with sub-diffraction-limited precision (Churchman et al. 2005), this virtually renders multi-color SMT in the living cell impossible, as in the living cell specific problems with aberrations and chromatic effects inherent to the cellular environment need to be overcome. This problem appears to be overcome by the recent development of a ‘super-registration’ technique for single molecules with high spatial precision between spectral channels. Using a nuclear pore fluorescent stain it was possible to achieve a 10-nm local registration of nuclear pores and endogenous mRNA, labeled with the MS2 system. “Super-registration”
allowed following interactions of single mRNAs and pores during export and resolving individual transient steps of the export process and their respective rate constants, which were previously undefined (Grunwald and Singer 2010).

In conclusion, the HIV life cycle demonstrates how important it is to gather a complete picture in time and space of cellular and viral mechanisms that, without microscopy, might not be possible. The horizon for the future should include single-molecule studies on the virus’ dynamic equilibrium of processes in living cells that, until now, have been out of reach. Even more, the perspective of investigating individual complexes and molecular interactions in the living cell at the single-molecule level, e.g., through developments like “super-registration”, guides the way towards a quantitative picture of the dynamic equilibrium of cellular life.

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