Partition of Myc into Immobile vs. Mobile Complexes within Nuclei

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Myc levels are highly regulated and usually low in vivo. Dimerized with Max, it regulates most expressed genes and so directly and indirectly controls most cellular processes. Intranuclear diffusion of a functional c-Myc-eGFP, expressed from its native locus in murine fibroblasts and 3T3 cells or by transient transfection, was monitored using Two Photon Fluorescence Correlation Spectroscopy, revealing concentration and size (mobility) of complexes. With increased c-Myc-eGFP, a very immobile pool saturates as a 'mobile' pool increases. Both pools diffuse too slowly to be free Myc-Max dimers. Following serum stimulation, eGFP-c-Myc accumulated in the presence of the proteasome inhibitor MG132. Stimulating without MG132, Myc peaked at 2.5 hrs, and at steady was ~8 ± 1.3 nM. Inhibiting Myc-Max dimerization by Max-knockdown or drug treatment increased the 'mobile' c-Myc pool size. These results indicate that Myc populates macromolecular complexes of widely heterogenous size and mobility in vivo.
subsequent autocorrelation analysis are: 1) $G_0$, the amplitude at delay time zero—simply related to the inverse of the number of particles diffusing in the small volume and 2) the translational diffusion time. To obtain the self diffusion coefficient, the autocorrelation function must be analyzed with a particular physical model such as 3D diffusion, anomalous diffusion or diffusion with binding.

Fluorescence Correlation Spectroscopy (FCS) exploits small fluctuations of concentrations by monitoring very small volumes. In fact, the smaller the concentration that can be observed, the larger the fluctuations that can be recorded and consequently the larger the autocorrelation signal. Hence the maximal correlation ($G_0$) is inversely proportional to the total concentration. For a protein expressed at the few nanomolar level, like c-Myc, FCS is one of the best methods available to monitor dynamics in vivo.

**Results**

To insure that c-Myc-eGFP was appropriately expressed and physiologically regulated, cells derived from a mouse homozygous for a c-myc allele fused in frame at the end of its coding sequence to DNA encoding a destabilized eGFP; the chimeric Myc-eGFP from these cells is fully functional and expressed at physiological levels, and turns over normally. Myc-eGFP was readily observed in MEFs by standard fluorescence microscopy only after serum-induction and stabilization using the proteasome inhibitor MG132 (supplementary Figure SN1). To reliably detect the low levels of c-Myc-eGFP within the MEF nuclei, we used a narrow eGFP emission filter coupled with two photon (2p) excitation in the red at 970 nm. We noticed that within the (-GFP) nucleus there was relatively little autofluorescence. As seen in Figure 1, a high resolution scan barely detects the c-Myc-eGFP in a MEF cell stimulated with serum after 30 minutes, even though this level ($\sim 10$ nM) is higher than the steady state level measured (8 ± 1.3 nM). Using FCS, we are able to detect c-Myc-eGFP at very low concentrations without causing visible photodamage to the cell. Figure 1C (in red) compares the autocorrelation traces of the c-Myc-eGFP +ve cells with those of a wild-type MEF lacking GFP. Note the long “tail” in the autocorrelation function (ACF) of the c-Myc-eGFP protein, indicative of a quite immobile pool. At 970 nm excitation, for cells lacking GFP, it is clear there is very little auto-fluorescence, and better yet, this component has no apparent correlation (Fig. 1C, black).

We found that the overall population of c-Myc-eGFP in these MEF cell nuclei was stable, and that it could be divided into two sub-populations with different diffusion coefficients. These two diffusion coefficients were extracted repeatedly from the ACFs of different cell nuclei. Statistically identifying and separating a third translation component is only possible when three very disparate mobilities are evident, and this was not the case in this data. We thus decided to identify only two (arguably mixed) fractions, one denoted “mobile” and one “less mobile”. This simplification was needed despite the fact that the “tail” can be very smooth at times, hinting at lesser contributions from multiple slow diffusion coefficients and/or other dynamic processes like transient binding. Figure 1D shows the ACFs for the two separate components with the appropriate $G_i(t)$’s; we separate them into a faster or more mobile fraction (green) and a less mobile or immobile (almost) fraction, blue.

**Figure 1 | MYC partitions into less and more mobile populations.** (a) MEF cell with c-Myc-eGFP excited at 800 nm. (b) Same cell excited at 970 nm (high res scan of 1 ms/pixel), this cell contains about 15 nM of c-Myc-eGFP by FCS. (c) Autocorrelation function of a MEF eGFP-cMyc cell excited at 970 nm (red) and a wildtype MEF with no clear autocorrelation (black). (d) An example of the two component fit performed for ACF is shown.
Figure 2a shows the apparent change in populations in three different cells manipulated to produce different concentrations of c-Myc-eGFP. These include transient transfection of eGFP-c-Myc into wild-type or into the homozygous MEFs. (It should be noted that expression of the transfected Myc over the short intervals required for these experiments was not associated with any obvious changes in cellular or subcellular phenotype.) The low levels of expression are exemplified in Figure 2B, where a 2p image of a cell containing 100 nM of c-Myc-eGFP is barely visible. The image was taken over a long exposure (at a 1 ms/pixel duration) and moderate resolution (256 × 256 pixels). As can be seen in Figure 2c, the more “mobile” fraction increases as the total expression of transfected protein increases; concurrently, the “immobile” fraction remains nearly constant. At 100 nM, the more mobile fraction represents >90% of the total c-Myc concentration.

We also studied the physiological accumulation of c-Myc-eGFP over time after serum stimulation of serum-starved MEFs. The addition of the drug MG132 was used in some cells to prevent the degradation of c-Myc-eGFP allowing us to follow the stimulated accumulation for 6–7 hrs. The results are shown in Figure 3A, and match the expected temporal pattern. There is a peak at 2–3 hrs for the cells without MG132 (with an increase of about 30% from the final steady state level of about 8 nM). For those with MG132, we see a prolonged and exaggerated increase of cMyc-eGFP concentration. Also, while the concentrations of both fractions increase, the more mobile fraction rapidly becomes dominant (data not shown). We also considered the expected noise for what is essentially an expression counting experiment. Figure 3B displays the coefficient of variation (CV), defined as the ratio of the standard deviation to the mean, for MG132 and nontreated cells. The CV spikes one hour after

**Figure 2** | Increased MYC augments mobile-MYC (a) ACF of three MEF cells with different concentrations of eGFP-cMyc. The concentrations are 10 (blue), 94 (red) and 400 nM (black). All three ACFs have been normalized to G(blue) at τ = 1e − 3. (b) Image of MEF cell at 970 excitation containing about 50 nM of eGFP-cMyc (high resolution). (c) The fractions of the mobile and immobile populations obtained from transfections of wild type and homozygous cells are shown.
addition of rich serum medium. The large increase is likely due to lack of synchrony in the cells in response to the impulse and consequent variation in nuclei of different cells. Despite accumulating considerably more Myc protein, the CV of the MG132 treated cells is slightly larger than for nontreated cells. CV for a counting dominated process should decrease by 1/square root of the controlling number; thus the relevant number in these nuclei (perhaps the number of active transcripts, see discussion) did not need to increase to increase c-MYC numbers.

The diffusion of even the most mobile fraction was considerably slower than would be expected for a freely diffusing Myc-eGFP-Max dimer. To test if the mobility of either the immobile or mobile fractions depended on DNA binding, cells at steady state were treated with the inhibitor of Myc-Max dimerization 10058-F4\textsuperscript{19}. Because structural studies have revealed that DNA-binding by Myc is stringently dependent on such dimerization\textsuperscript{20}, this inhibitor was expected to distinguish DNA-binding dependent and independent subpopulations of Myc. This same inhibitor has been demonstrated to reduce Myc binding at target promoters genome-wide, in vivo\textsuperscript{10}. Figure 4 shows the distribution of diffusion coefficients of the recovered 'mobile' fraction. The mean Dt of the mobile fraction for these cells

Figure 3 | Its coefficient of variation indicates that MYC is regulated at several steps. (a) The c-Myc concentration of MEF cells treated and non-treated with MG132 over a period of 6–7 hrs. Left axis is the concentration of the MG132 treated cells (RED) and right axis is concentration of non-treated cells (BLACK). (b) Coefficient of Variation (CV) for nontreated and MG132 treated cells.
was nearly unchanged: 3.4 ± 1.6 and 3.0 ± 1.8 μm²/sec with and without inhibitor (10058-F4) respectively. Those broad error limits represent biological variation, however, and it is clear upon inspection that the mobile fraction histogram skewed towards a more mobile distribution when the inhibitor was added. Although 10058-F4 shifted c-Myc-eGFP to slightly higher mobilities, it did not provoke the wholesale release of Myc to freely diffusing forms; thus, it seems that even without Max, Myc remains associated with its

Figure 4 | Inhibition of MYC-Max dimerization does not liberate freely diffusing MYC. (a) Diffusion coefficient shifted slightly with the addition of inhibitor. (n = 42 for no inhibitor and 37 for inhibitor). (b) Max knockdown cells have a slightly larger diffusion coefficient to start and the inhibitor further increases this fraction (n = 24 for no inhibitor and 32 for inhibitor).
other macromolecular partners. At very high concentrations of transfected c-Myc-eGFP, a diffusion coefficient of 5.5 ± 0.3 μm²/sec was recovered for the most mobile fraction, suggesting that MYC may eventually saturate those lower mobility partners and “spill over” into a somewhat higher mobility niche. We considered using a ’global’ approach for the analysis of the diffusion coefficients, but at this stage we were more interested in obtaining data on Dt variations in many cells and the qualitative patterns that could be discerned. Eventually, when we return to the mosaic nature of expression, global analysis will become a key element in quantifying populations and rates.

RNAi or shRNA that targeted Max efficiently (supplementary Fig. SN2) were also used in combination (or not) with 10058-F4 to liberate Myc from Myc-Max dimers in cells at steady state. While these manipulations failed to expose a new highly mobile population of “free” Myc-eGFP, the diffusion times recovered from these measurements indicated noticeable increases in the mobile fraction. The Dt of the immobile fraction remained near 0.20 ± 0.06 μm²/sec with all

Figure 5 | Without Max, MYC complexes are more mobile. (a) Mobile fraction shifted slightly with the addition of inhibitor (n = 42 for no inhibitor and 57 for inhibitor). (b) Knockdown cells have a slightly larger population of mobile fraction to start and the inhibitor further increases this fraction (n = 24 for no inhibitor and 32 for inhibitor).
levels will, of course, require additional experiments that manipulate exactly why CV is elevated above protein copy number – predicted generating (on average) about 10 c-MYC. Precise determination of the length of the experiment. The difference could be thought of as a about 3%, while our increased CV lies within the range 10–20% over error expected for a protein with copy number near 1000 is only cell-to-cell variation in c-Myc levels in MG132-treated cells would tion of c-Myc-eGFP rising to 400% of the steady state levels. On hours post-stimulation; treatment with the proteasome inhibitor transfection that at c-MYC levels beyond physiological, distinctly work within the nucleus. Upon flooding the cell with large quantities of exogenously expressed c-Myc-eGFP, the excess protein partitions into the nucleus, despite low copy numbers, points toward a future where this technique is employed to follow the dynamics of transcriptional activation pro cesses throughout the cell cycle and to assess the possibly functional heterogeneity within a population of cells. Few methods can directly monitor concentration of a key protein at nM levels; fewer still can do so nondestructively (allowing sequential observation throughout cell cycle or experiment condition changes) and even fewer can explore the mosaic nature of expression via histograms. FCS is thus a desirable technique for studies of regulators.

Methods

2P FCS. Two photon imaging and FCS measurements were carried out using a system built in the Ultrafast Laser Microscopy laboratory. The excitation source was a tunable Ti-Sapphire broadband Mai Tai laser (Spectra-Physics-Newport) set at 970 nm. The excitation power was set at a level where no bleaching or visible damage to the cell occurred (~10 mW at the microscope entrance). The microscope was a Zeiss Axiosvert 135 M using an E680 SP 2P dichroic filter (Chroma Technology Corporation) to eliminate the IR exciting light. The objective was a 100× Plan Neofluar oil objective (Zeiss) with NA 1.3. The microscope was equipped with a piezo-electric stage for x-y control (Mad City Labs.) and the objective was also equipped with a piezo-electric device for z-control (Mad City Labs). Bone marrow was used an Alba 3 channel fluorescence correlation system (ISS, Inc.). The detected light was split into the two channels with a 495 nm dichroic mirror (Chroma). A 515/40 nm bandpass emission filter was placed before channel 1 for eGFP detection, and a 450/40 nm bandpass filter was used with channel 2 to detect autofluorescence. Two photon volumes were calibrated using a transgenic mouse strain of C57BL/6J that expresses fluorescent beads. The bead beam waist in plane, w0, was found to be 0.34 μm, while the axial extent zw was 1.8 μm. The diffusion coefficient of GFP in a cell has been measured to be 26 ± 7 μm2/s. Also, a mutant of the transformation factor VBP that does not bind DNA and has only a “leucine zipper” domain was measured to have a diffusion coefficient of 13 ± 4 μm2/s. These values are in agreement with the diffusion coefficients of NLS-eGFP proteins.

Cell culture and transfections. In some experiments, a plasmid vector transiently directing the expression of the same c-Myc-eGFP fusion was transfected into these same homozygous or non-targeted (wild-type) fibroblasts. Cells were incubated with the inhibitor 10058-F4 for at least six hours. Knockdown treated cells were incubated for 48 hours before measurements.

Preparation of MEF and 3T3 cells. MEF cells were prepared following standard protocols (http://www.fhcrc.org/science/lab/fibro/protocols/MEF.html). In brief, E13.5 embryos were harvested from wt or homozygous c-Myc-eGFP knock-in mice and washed with sterile PBS to remove any remaining maternal blood cells. After removal of the embryonic cord, liver, spleen and tubular intestine, the bulk of the CNS tissue was dissected by dissection the head above the level of the oral cavity. The remaining embryo was treated with trypsin (5 ml/embryo) and minced with sterile forceps and scissors for 2–3 minutes and incubated in 37°C for 15 minutes. The embryonic tissue of embryo was further dissociated by triturating with a 10 ml pipet. Isolated primary MEF cells were pelleted in 5 ml of DMEM with 10% FBS by centrifugation at 1000 RPM for 5 min. After aspirating the supernatant, the P0 primary MEF cells were re-suspended in fresh medium and incubated at 37°C in 5% CO2. The 3T3 cells were cultured on a 1.2 × 10^5 primary MEF cells every 3 days on PDL dishes (http://labs.fhcrc.org/fibro/Protocols/MEFS.html; Nilausen K. Green H. Exp Cell res. 1965). After a rapid growth period (passage 1–5), a slower growth period (50–10) and senescence (little or no growth, passage 10–25) the immortalized 3T3 grow out.

Serum starvation and stimulation. The WT, c-Myc-eGFP primary and/or NIH 3T3 cells were cultured in DMEM containing 0.03% FBS for 48 hrs, then trisipnded and re-cultured in the DMEM medium containing 10% FBS and 1% MEM-Non-Essential Amino Acids (MEM NEAA, GibCO 11140) in the presence and/or absence of proteasome inhibitor MG132 or Myc-MAX inhibitor, 10058-F4. The cells were seeded in the Lab-tek Chambered Coverglass w/cv #1German borseillary tic two chambers slides (NUNC cat. # 155380) and incubated at 37°C in 5% CO2. Pheno-less DMEM (GibCO, 20163) and charcoal/dextran treated FBS (HyClone, SH30068.01) were used during analysis.

Knock-down MAX gene. Custom designed stealth siRNA for MAX accession No. BC138617 position 313, 5′-CCAUAGCCACCCCAAGAACAUUGG-3′ and control, 5′-AUAUGCUUGCUUGCGCGUAUGG-3′ (Invitrogen) were electroporated to either primary or NIH 3T3 cells with Amaxa MEF2 Nucleofector Kit (LONZA, VPD-1005) using program MEF2/A023 followed manufacturer’s protocols. About 2.5 mmole of each stealth siRNA were used for one well of a 6 well plate and incubated at 37°C, in 5% of CO2 for 40–48 hrs for assay.

The ability of FCS to nondestructively report on both mobile and chromatin-associated populations of c-Myc in nuclei, despite low copy numbers, points toward a future where this technique is employed to follow the dynamics of transcriptional activation processes throughout the cell cycle and to assess the possibly functional heterogeneity within a population of cells. Few methods can directly monitor concentration of a key protein at nM levels; fewer still can do so nondestructively (allowing sequential observation throughout cell cycle or experiment condition changes) and even fewer can explore the mosaic nature of expression via histograms. FCS is thus a desirable technique for studies of regulators.
MAX knock-down stable cell line. PLKO.1 lentiviral vector carrying MAX shRNA/ RNAs sequences 5’-AAAGCTGTCTTTGATGTTGTC-3’ (Thermo-Open Biosystem, cat. no. RHS979-9607256, clone ID: TRCN0000039867) was transfected into HEK 293T cells with arrest-in reagent according to manufacturer’s protocol. Viral particles were packaged, titered and transduced into the c-Myc-eGFP NIH 3T3 cells to create the MAX stable knock-down cell line. The PLKO.1 vector plasmid was the negative controls for these assays.

Q-real-time PCR analysis. Total RNAs were purified from WT, c-Myc-eGFP and c-Myc-eGFP-MAX knock-down NIH 3T3 cells at various time points as described (Nie et al., 2012). 0.25 mg of total RNA from each time point were used for first-strand c-DNA synthesis (Enhanced Avian HS RT-PCR-100 kit, SIGMA, Cat. No. HSRT100-1kt). The primers and probes for each gene were designed using Roche Universal Probe library Assay Design Center Web (Figure S7F). q-PCR was performed with the Roche LightCycler 480 system (LightCycler 480 Probe Master, Ref. No. 04 707 494 001; Universal Probe library set, Human, Ref. No. 04 683 653 001). The expression levels of c-Myc, Max and MAD were normalized with GAPDH.

Immunoblotting. Equal amounts of c-Myc-eGFP-MEF and MAX knock-down cells at different time points were harvested in RIPA buffer (150 mM NaCl, 1% NP-40, 0.5% NaDOC, 0.05% SDS, 50 mM Tris, pH 7.5 and 1× protease inhibitor). Proteins were separated on 4–12% SDS-PAGE and blotted with anti-c-Myc (Epitomics, Cal. No. Y69), anti-Max (C-17) (Santa Cruz Biotech, Cal. No. SC-197), anti-Mad 1 (FL-221) (Santa Cruz Biotech, Cal. No. SC-766) and anti actin (1-19) (Santa Cruz Biotech, Cal. No. SC-1616).

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