Novel Tools and Methods

Heterogeneities in Axonal Structure and Transporter Distribution Lower Dopamine Reuptake Efficiency

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Abstract

Efficient clearance of dopamine (DA) from the synapse is key to regulating dopaminergic signaling. This role is fulfilled by DA transporters (DATs). Recent advances in the structural characterization of DAT from Drosophila (dDAT) and in high-resolution imaging of DA neurons and the distribution of DATs in living cells now permit us to gain a mechanistic understanding of DA reuptake events in silico. Using electron microscopy images and immunofluorescence of transgenic knock-in mouse brains that express hemagglutinin-tagged DAT in DA neurons, we reconstructed a realistic environment for MCell simulations of DA reuptake, wherein the identity, population and kinetics of homology-modeled human DAT (hDAT) substates were derived from molecular simulations. The complex morphology of axon terminals near active zones was observed to give rise to large variations in DA reuptake efficiency, and thereby in extracellular DA density. Comparison of the effect of different firing patterns showed that phasic firing would increase the probability of reaching local DA levels sufficiently high to activate low-affinity DA receptors, mainly owing to high DA levels transiently attained during the burst phase. The experimentally observed nonuniform surface distribution of DATs emerged as a major modulator of DA signaling: reuptake was slower, and the peaks/width of transient DA levels were sharper/wider under nonuniform distribution of DATs, compared with uniform. Overall, the study highlights the importance of accurate descriptions of extrasynaptic morphology, DAT distribution, and conformational kinetics for quantitative evaluation of dopaminergic transmission and for providing deeper understanding of the mechanisms that regulate DA transmission.

Key words: Dopamine reuptake; dopamine transporter; spatial simulation; stochastics

Significance Statement

Dopamine (DA) modulates motor control, cognition, and drug addiction. Understanding the mechanism of dopamine transmission is essential to designing therapies for neurologic disorders. We developed a multiscale model using advances in imaging and high-performance computing technologies, which permitted us to perform spatially realistic simulations of dopamine reuptake. Simulations show large temporal and spatial variations in the local density of dopamine depending on the morphology of the synaptic/extrasynaptic region near dopamine release site and on firing pattern. Dopamine clearance is less efficient under heterogeneous distribution of dopamine reuptake transporters, compared with a uniform distribution with the same average surface density. Dopamine reuptake transporters membrane distribution, accessibility of dopamine reuptake transporters outward-facing conformation, and large fluctuations in dopamine levels emerge as key features that modulate dopaminergic transmission.

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Introduction

Midbrain dopaminergic neurons have a strong influence on striatum functions such as motor and action planning, cognitive functions, and motivation (Roepner, 2013). Dys-regulation of dopaminergic transmission leads to impairment of these activities, resulting in disorders such as Parkinson’s disease (PD; Hoang, 2014), attention deficit/hyperactivity disorder (ADHD; Wu et al., 2015), and drug addiction (Nutt et al., 2015). A mechanistic understanding of dopamine (DA) transmission events is essential to developing therapeutic strategies, because many behavioral states strongly correlate with DA release and/or reuptake (Tsai et al., 2009; Sulzer et al., 2016).

DA release to the synapse is activated by excitatory stimulation and exhibits patterns similar to neuronal firings. DA excitatory signaling proceeds by activation of DA receptors on binding DA molecules. DA rapidly diffuses from the active zone (AZ; or release site) to extrasynaptic regions in the extracellular (EC) medium. DA transporters (DATs), membrane proteins usually located on the surface of presynaptic axon terminals, regulate DA signaling by removing excess DA from extrasynaptic regions (Torres et al., 2003; Vaughan and Foster, 2013). DATs are targets for addictive substances, which inhibit their function (Amara and Sonders, 1998), thus resulting in excess (neurotoxic) DA levels in the EC region (Di and Imperato, 1988), whereas low levels of DA cause motor impairments associated with PD (Lotharius and Brundin, 2002). In addition, reuptake by DATs, the rate of DA diffusion from the AZ to the extrasynaptic region (Taylor et al., 2013), and the frequency and patterns of action potentials (APs; Tsai et al., 2009) are known to modulate the efficiency of DA signaling.

The dynamics of DA reuptake by DATs has been a focal topic in modeling efforts, at both the cellular and molecular levels (Mortensen and Amara, 2003). Early efforts at the cellular level adopted a well-mixed model focusing on predicting the DA concentration at certain regions of the brain, such as the nucleus accumbens or dorsal striatum (Garris et al., 1994). The effect of EC DA concentration on the activation of DA receptors (Viggiano et al., 2004), as well as spatial/volume exclusion/transmission properties affecting EC DA levels, have been included in later volume transmission (VT) models (Cragg et al., 2001; Cragg and Rice, 2004; Rice and Cragg, 2008; Dreyer et al., 2010; Dreyer and Hougsaard, 2013; Sulzer et al., 2016). These studies highlighted a need for considering the distinctive diffusion and uptake characteristics of the EC microenvironment. Yet, no quantitative models/simulations have been developed or performed to date that would permit us to assess how the complex geometry of DA terminals and the spatial distribution and conformational dynamics of DATs alter dopaminergic signaling. Advances in imaging DA neurons and visualizing individual DATs (Block et al., 2015) now enable us to reconstruct in silico the detailed morphology near AZs and examine the time evolution of DA release and reuptake with the help of MCell, software originally developed (Stiles et al., 1996; Kerr et al., 2008; Czech et al., 2009) for spatiotemporally realistic simulations of synaptic signaling events.

In addition to cellular structure and heterogeneities, the conformational dynamics of DATs is a determinant of DA transport efficiency. Recent crystal structures of Drosophila DAT (dDAT; Penmatsa et al., 2013, 2015; Wang et al., 2015) have opened the way to structure-based studies of DAT dynamics. Simulations based on these structures helped elucidate the sequence of molecular events that take place during the transport cycle of the human orthologue, hDAT (Cheng and Bahar, 2015; Khelashvili et al., 2015; Razavi et al., 2017; Cheng et al., 2018). We are now able to make reasonable approximations for the kinetic scheme and parameters associated with the DAT transport cycle based on statistical analyses of the full-atomic trajectories and free energy calculations.

Here, we present an integrated model of synaptic signaling in DA neurons developed from cellular and molecular structures and molecular dynamics. We investigate the effects of (1) the conformational kinetics of DATs, (2) the spatial complexity of DA terminals and AZs based on fluorescence images, (3) the firing patterns, phasic versus tonic, and (4) the heterogeneous distribution of DATs on the plasma membrane based on experimentally observed DAT density fluctuations. Simulations reveal the strong dependence of local DA levels as well as overall DA clearance efficiency on the local geometry of axon terminals. They also reveal that the presence of DAT clusters (consistent with the DAT density heterogeneities observed in high-resolution images; Block et al., 2015) causes a reduction in the efficiency of DA reuptake compared with uniformly distributed DATs with the same average surface density. This effect becomes more pronounced with increasing heterogeneity of the surface distribution of DATs.
Materials and Methods

Confocal imaging of immunolabeled DATs in transgenic mouse brains

The procedure for preparing and imaging acute brain slices from transgenic knock-in mice of either sex expressing DAT molecules tagged with the hemagglutinin-A (HA) epitope (HA-DAT; (Rao et al., 2012) has been described in previous works (Rao et al., 2013; Block et al., 2015). Briefly, brains were submerged into an ice slush of oxygenated artificial cerebrospinal fluid (ACSF), and 0.8-mm-thick sagittal slices were cut using microtome blades and a stainless-steel slicing block. The subcellular localization of cell-surface HA-DAT molecules was deduced from intact living DA neurons in acute sagittal brain slices as detected by mouse anti-HA antibodies with Cy3-conjugated anti-mouse antibodies (Block et al., 2015). Slices were incubated in ACSF at room temperature with 1 μg/ml mouse anti-HA antibodies for 1 h. After removing unbound antibodies, slices were incubated for 1 h at 4°C in ACSF with 2.5 μg/ml Cy3-conjugated anti-mouse Fab fragments.

We have not observed substantial differences in HA-DAT distribution between live-cell and postfixation labeling with HA antibodies, suggesting that axonal varicosities revealed by DAT staining were not the result of blebbing during the labeling procedure of live slices. We found that binding of antibodies to live neurons followed by fixation provides much superior image quality and lower signal-to-noise ratio compared with the conventional protocol of fixation first and then staining with antibodies. Importantly, live-neuron staining protocol allows labeling of cell-surface DATs, which is important for defining the distribution of DATs on the neuronal surfaces in the model. Observations of DAT endocytosis (Block et al., 2015), normal lateral membrane mobility of DAT, and healthy mitochondria in dopaminergic neurons from slices kept alive for at least 2 h were indicative of functional neurons in these slices. Moreover, DA neurons labeled as described above have been observed to exhibit pH-dependent vesicular trapping of antipsychotic drugs (Tucker et al., 2015).

To obtain high-resolution 3D images of DA neurons, a z-stack of 18 confocal images at 400-nm interstack distance was acquired 10 μm deep from the cut face of the slice through the 561 filter channel using a spinning disk confocal system based on a Zeiss Axio Observer Z1 inverted fluorescence microscope (with 63× Plan Apo PH NA 1.4 objective), equipped with a computer-controlled Spherical Aberration Correction unit, Yokogawa CSU-X1, Vector photo manipulation module, Photometrics Evolve 16-bit EMCCD camera, Hamamatsu CMOS camera, environmental chamber, and piezo stage controller and lasers (405, 445, 488, 515, 561, and 640 nm), all controlled by SlideBook 6 software (Intelligent Imaging Innovation).

Our image reconstruction and modeling described below are based on the combination of light microscopy images of slices and electron microscopy images (Block et al., 2015) that were obtained on intact animals after cardiopertusion fixation.

In silico reconstruction of DA axonal terminals in the striatum

We reconstructed in silico a 10 × 10 × 7.2-μm volume from the above described striatal region (Fig. 1A–C) using a semiautomated 3D reconstruction algorithm (Turetken et al., 2016). The size of the simulation box was large enough to allow for diffusion of DA over a sufficiently broad EC region, in accord with previous estimates (Ven-ton et al., 2003), and the reconstruction yielded a realistic representation of both the heterogeneous shape of axonal terminals and the surface distribution of individual DAT molecules. The reconstructed region contained 13 axon terminals (Fig. 1D). The corresponding volumes and surface areas (listed in Table 1) were calculated using the NeuroMorph (Jorstad et al., 2015), a Blender add-on that uses triangular meshes to evaluate the surface area and corresponding normal to determine the volume of each tetrahedron. The total volume occupied by the 13 axon terminals was 101.03 μm3, and the corresponding total surface area, 337.16 μm2.

The DA axonal terminals reconstructed in silico contained six varicosities, i.e., 3D globular regions with densely expressed DATs, distributed over three DA terminals: one of the largest terminals had three varicosities, another had two, and the remaining varicosity was on a third terminal. AZs lie within varicosities but are not usually populated with DAT molecules (Block et al., 2015); accordingly, subregions (of varicosities) that lacked DAT molecules within at least a 50-nm radius were identified as AZs. The region between DAT-expressing cells (detected by fluorescence microscopy) and others (not visible) was represented by an interstitial (void) space of 30-nm thickness surrounding the DAT-expressing terminals (Fig. 1E). This led to a void fraction of 0.21, consistent with previous estimates (Cragg et al., 2001), or an overall volume of 155.35 μm3 (Table 1) that formed available for DA diffusion. These narrow regions form the synaptic clefts and extrasynaptic regions available for the diffusion of DA molecules. The number of AZs for a given volume was verified to be comparable to that used in other studies (Dreyer et al., 2010).

Next, we describe the placement of DAT molecules on the membrane of axonal terminals. To investigate the effect of DAT surface distribution heterogeneities on the efficiency of DA reuptake, we examined four cases (Fig. 2). Case 1 refers to the uniform distribution (Fig. 2A), taken as μ(DAT) = 800/μm2, based on the electron microscopy images of gold particle labeled HA-DAT, assuming 10% labeling efficiency (Block et al., 2015). Case 2 is a non-uniform (bimodal) distribution (Table 2; Fig. 2B), set forth in accord with the actual distribution of DATs observed in experiments. High-density regions were detected in the fluorescence images as continuous bright regions (Fig. 1C). These regions covered ~10% of the plasma membrane area, and ~90% of DATs were localized in these regions. The surface densities of DATs in the high- and low-density regions were taken as ρh = 6339/μm2 and ρl = 50/μm2, respectively. In case 3, the distribution is again bimodal, similar to case 2, but the central parts of the high-density regions from case 2 are selected as the
new very-high-density regions, with $\rho_h = 30,000/\mu m^2$ and $\rho_l = 50/\mu m^2$ elsewhere, which leads to a sharper heterogeneity in the spatial distribution of DATs (Fig. 2C).

In case 4, DATs are assumed to be clustered in the immediate neighborhood of AZs, as a mimic for conventional synaptic models where DATs act as gatekeepers near the synaptic cleft (Fig. 2D; Rothstein et al., 1994; Danbolt, 2001), and DAT surface concentrations in high- and low-density regions are the same as in case 3. The histograms in Fig. 2E–H describe the probability distribution of the distances of DAT molecules from the closest AZ.

Table 1. Geometric characteristics of axon terminals reconstructed\(^1\) for simulations

| Subsystem V (\(\mu m\))^3 | Surface area (\(\mu m^2\))^2 | Available\(^3\) surface area (\(\mu m^2\))^2 |
|---------------------------|-----------------------------|------------------------------------------|
| DA axons                  |                             |                                          |
| Terminal 1                | 1.22                        | 6.8                                      |
| Terminal 2                | 24.63                       | 59.47                                    |
| Terminal 3                | 3.53                        | 20.18                                    |
| Terminal 4                | 15.84                       | 56.33                                    |
| Terminal 5                | 4.72                        | 25.14                                    |
| Terminal 6                | 1.71                        | 10.12                                    |
| Terminal 7                | 0.82                        | 5.57                                     |
| Terminal 8                | 2.04                        | 12.94                                    |
| Terminal 9                | 15.32                       | 46.86                                    |
| Terminal 10               | 15.18                       | 58.65                                    |
| Terminal 11               | 3.85                        | 18.99                                    |
| Terminal 12               | 9.69                        | 34.31                                    |
| Terminal 13               | 2.58                        | 12.35                                    |
| Total for all terminals   | 101.03                      | 367.71                                   |
| Non–DA-expressing cells   | 463.62                      | 1058.00                                  |
| DA axons + other cells    | 564.65                      | 1425.71                                  |
| Available EC volume\(^2\) | 155.35                      |                                          |

\(^1\) Using experimental data from Block et al. (2015).

\(^2\) Obtained by subtracting 564.65 from 720 \(\mu m^3\) (for the simulation box of \(10 \times 10 \times 7.2 \mu m^3\)).

\(^3\) Excluding those at the simulation box boundaries.

MCCell simulations of DA release and reuptake events in DA neurons

Spatiotemporally realistic simulations were performed using MCell (Stiles et al., 1996; Bartol and Stiles, 2001; Kerr et al., 2008), a 3D reaction-diffusion system solver that allows users to reconstruct complex geometries, define the subcellular localization of discrete molecules, and simulate their dynamics. The parameters used in simulations are given in Table 2. Unimolecular reactions are scheduled according to defined reaction rates, and bimolecular reactions occur with predefined probabilities that are chosen to match bulk reaction rates. Collisions between molecules are detected by ray-tracing algorithm. We adopted Neumann boundary conditions similar to
those used in recent simulations of Ca\textsuperscript{2+} signaling (Bartol et al., 2015c), i.e., DA molecules are subjected to reflective boundary conditions at the simulation box walls. In addition, to reduce the bias from reflective boundaries, terminals within 1 \textmu m of the box boundary were assumed to be inactive, such that the available surface area on DA axon terminals was 337.16 \textmu m\textsuperscript{2} (see Table 1).

The probability of a release succeeding an action potential depends on multiple factors (Dreyer et al., 2010), including the content of DA in the striatum ($I_{DA}$; Bannon et al., 1981), the ratio of the amount of total DA released per action potential ($R$; Gubernator et al., 2009), the volumetric density of DA terminals at AZs ($\rho_{term}$; Doucet et al., 1986), and the number of DA molecules released per quantum ($N_0$; Pothos et al., 1998). The parameters are given in Table 2, which yielded a release event probability of 6\% (Dreyer et al., 2010). Each AZ has a release site located at its center; and on a release event, a total of $N_0$ DA molecules is assumed to be released from the release site. DA diffusion was modeled as a pseudorandom walk with a fixed time step of $\Delta t = 0.1 \mu s$. The distribution of DA step sizes yielded an average of 13.3 nm using a DA diffusion coefficient of $4 \times 10^{6}$ cm\textsuperscript{2}/s. A time step of 100 \mu s was used for the slow events, such as the transition of

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**Figure 2.** Four space-dependent models of different plasma distributions of DATs. A–D, Distant (left) and magnified (right) view of the axon terminals in cases 1–4. Each color shows a different terminal, and the red dots represent the DATs. The white regions in A represent the regions with high fluorescence intensity, and those regions are filled with high density of DATs (red dots), magnified in B, where the blue region shows the AZ. The red patches in B–D illustrate high-density regions where ~90\% of DATs are clustered. E–H, Distance distribution of DATs to closest AZ center, from 140 independent simulations.
DAT to reuptake-ready (EC-exposed outward-facing) state release of its cargo. 140 independent runs, each of 10 s, were performed to extract statistically significant results.

**Conformational dynamics of DAT**

Our recent dual-boost accelerated molecular dynamics (aMD; Hamelberg et al., 2007; Miao et al., 2013) and conventional MD (cMD) simulations of DAT dynamics showed that the DA transport cycle by DAT can be approximated by four basic steps (Fig. 3; Cheng and Bahar, 2015; Cheng et al., 2018): (1) recognition and binding of DA (and cotransported Na\(^+\) ions) from the EC region to DAT in the outward-facing (OF) state—we designate the substrate- and Na\(^+\)-bound (or loaded) OF state as OF\(^+\); (2) global structural change of DAT from OF\(^+\) to inward-facing loaded (IF\(^+\)) state; (3) release of cargo to the IC region (IF\(^+\) → IF); and (4) reverse transition of the unbound/apo DAT from IF to OF state. The respective forward rate constants are denoted as \(k_{12}\), \(k_{23}\), \(k_{34}\), and \(k_{41}\), and reverse rate constants are \(k_{21}\), \(k_{32}\), \(k_{43}\), and \(k_{14}\) (Fig. 3).

The molecular events of DA binding and unbinding to DAT generally involve local conformational changes, and their energetics can be estimated using established free energy calculation methods. We used two methods: (1) alchemical free energy calculation with free energy perturbation (FEP) method (Pohorille et al., 2010) and (2) potential of mean force (PMF) calculations using the adaptive biasing force (ABF) method (Chipot and Hénin, 2005), based on cMD trajectories. The FEP calculations yielded a binding free energy change of \(\Delta G_{\text{bind}} = -7.8\) kcal/mol (Cheng et al., 2018), in excellent agreement with the experimental value of \(-7.4\) kcal/mol (Dar et al., 2006; Huang and Zhan, 2007). MD simulations indicated that the average time required to bind a DA molecule originally placed at a distance of 15 Å from the binding site is \(\sim 125\) ns. To convert this number into the binding rate constant \(k_{12}\), we performed the following operation. First, using an EC DA concentration of 7.5 nM (Feifel et al., 2003), we calculated the density of DA molecules to be \(7.5 \times 10^{-9}\) mol/nm\(^3\) \(\times 6.02 \times 10^{23}\) molecules/mol = 4.5 \(\times 10^{-9}\) molecules/nm\(^3\). The free volume (excluding that occupied by DAT itself) for DA translocation originally located at a separation of 15 Å from the binding was evaluated to be 2 nm\(^3\) using POVM (Durrant et al., 2011). The number of DA molecules colliding with DAT based on this accessible volume is 2 nm\(^3\) \(\times 4.5 \times 10^{-9}\) = 9 \(\times 10^{-6}\), which also represents the a priori probability/frequency of collision of a given DA molecule. This leads to an effective binding time of 125 ns/9 \(\times 10^{-9}\) = 13.88 s. By normalizing with respect to EC DA concentration, the bimolecular reaction constant is determined as \((1/13.88\) s)/7.5 \(\times 10^{-9}\) = 9.6 \(\times 10^{6}\) M\(^{-1}\) s\(^{-1}\).

We further observed that (1) the binding of Na\(^+\) ions was fast (<100 ns) and the subsequent binding of DA readily prompted the closure of the EC gate such that the escape of DA (and ions) back to the EC region was negligibly small, i.e., \(k_{21} \ll k_{12}\); (2) no DA efflux to EC region was detected (i.e., \(k_{43} = k_{32} \approx 0\)); (3) the DA-free (with Na\(^+\)/Cl\(^-\) bound) OF→IF transition \((k_{14})\) was even slower than the OF→IF transition \((k_{14})\) owing to the difficulty in closing the intracellular gates (Cheng et al., 2018). The global OF↔IF transition rates were thoroughly examined in microsecond aMD simulations of DA-free DAT, which showed that the population of reuptake-ready (OF) conformers was lower than that of IF (or other intermediate) conformers by a factor or 4, or \(k_{14}/k_{41} \approx 4\) (Cheng et al., 2018). These considerations provided us with robust information on the relative rates of the individual steps and led to the rate constants in Table 2, the absolute values of which were verified to be compatible with experimentally observed turnover rates and steady-state concentrations of DA molecules.

To investigate the sensitivity of DA reuptake efficiency to DAT conformational kinetics, we also performed global sensitivity analysis with respect to rate constants in Fig. 3. We performed 729 independent runs with different combinations of \(k_{12}\), \(k_{23}\), \(k_{34}\), \(k_{41}\), and \(k_{14}\) which we varied by three orders of magnitude. The results are presented in Fig. 4. Each blue dot represents the outcome, EC DA concentration in the simulation box, \([\text{DA}]_{\text{EC}}\), from one run. A broad range of \([\text{DA}]_{\text{EC}}\) values, from 0.1 to >100 nM, are observed, yet an increase in DA binding rate \(k_{12}\) results in a more efficient clearance and thereby lower DA levels in the EC region (Fig. 4A). A similar trend is detected with an increase in the transition rate \(k_{41}\) from IF to OF, which
exposes more reuptake-ready DATs to the EC region (Fig. 4C), and the reverse transition induces the opposite effect (Fig. 4D). An even sharper effect is observed on plotting \([\text{DA}]_{\text{EC}}\) against the ratio \(k_{14}/k_{41}\), highlighting the importance of the equilibrium population of the OF and IF states of DAT after releasing its cargo (Fig. 4E). The examination of the relative effects of DA binding \((k_{12})\) versus back-transition to the IF state \((k_{14}, \text{IF})\) for the OF DAT also indicates that the OF DAT level is a major determinant of \([\text{DA}]_{\text{EC}}\) (Fig. 4F). Further quantitative assessment of the statistical significance of these observations using Spearman rank correlation coefficients confirmed that the binding rate constant \(k_{12}\) and the ratio \(k_{14}/k_{41}\) are two major determinants of DA clearance efficiency. No clear effect was seen for \(k_{23}\) (Fig. 4B) or \(k_{34}\) (data not shown).

**Code accessibility**

The MCell software and the input files for the spatial models implemented in MCell are freely available to the community. The model files developed in this study can be accessed as Extended Data 1 or GitHub repository at [https://github.com/cihankayacihan/dopamine_striatum_mcell](https://github.com/cihankayacihan/dopamine_striatum_mcell). The only requirement for repeating the simulations is to download the MCell software ([http://www.mcell.org](http://www.mcell.org)) version 3.4 and use it together with the model files. The available files include information on geometry, molecules, reactions, surface classes, and release patterns in addition to output trajectories and MCell simulation parameters.

**Results**

*In silico* turnover and \([\text{DA}]_{\text{EC}}\) at half-maximal rate conform to experimental data

We first verified that MCell simulations yielded macroscopic properties consistent with experimental data. We calculated the turnover rate by adopting in our simulations the same protocol as that adopted in experiments (Rao et al., 2013): multiple runs are performed for a series of initial concentration of DA in the EC region, \([\text{DA}]_{\text{EC}}\), and in each case, the mass of DA molecules transported per unit time is measured. The number of DA molecules transported per second, \(V_{\text{max}}\), under saturation conditions (\([\text{DA}]_{\text{sat}}\) is of the order of tens of micromoles per liter) is used to evaluate the turnover rate as the ratio of \(V_{\text{max}}\) to the total number \(B_{\text{max}}\) of DAT molecules present in the

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*Figure 3.* Schematic representation of the sequence of events occurring during the transport cycle of DAT. A succession of four major states is observed in MD simulations: unbound and DA/ion-bound outward-facing states (OF and OF\(^*\)) followed by unbound and bound inward-facing states (IF and IF\(^*\)). The corresponding hydration patterns (water molecules in white and pink spheres) and interactions of intra- and extracellular gating residues (R85-D476 and R60-D436, respectively, in stick representation) are displayed. Green arrows indicate the transitions that were observed and evaluated in molecular simulations (see Table 2). The events indicated by the red arrows were unlikely \((k_{21})\) or not observed \((k_{32} \text{ and } k_{43})\) in MD runs. Curved arrows refer to the binding or unbinding of DA (purple, space filling), cotransported Na\(^+\) ions Na1 and Na2 (yellow spheres), and the chloride ion (blue sphere).
system. In our simulation environment, $B_{\text{max}} \approx 220,000$, based on fluorescence microscopy data (Block et al., 2015). To evaluate the turnover rate \textit{in silico}, we counted the number of DAs transported as a function of $[\text{DA}]_0$ and examined for each concentration the number of DAs translocated per second. This led to a reuptake rate of $1.2 \times 10^5$ DAs/s at saturation ($V_{\text{max}}$). Division by $B_{\text{max}}$ gave a turnover rate of 0.55/s, which is comparable to the reported values of 0.2/s (Rao et al., 2013), 0.9/s (Prasad and Amara, 2001), and 1.8/s (Prasad and Amara, 2001; Beuming et al., 2008; Rao et al., 2013).

The average DA level in the EC medium, $[\text{DA}]_{\text{EC}}$, observed \textit{in silico} after reaching steady-state conditions was 7.8 nM (Fig. 5A). The physiologic concentration of DA in the striatum varies between 5 and 50 nM (Owesson-White et al., 2012), consistent with the large fluctuations (of the order of $[\text{DA}]_{\text{EC}} \approx \pm 10$ nM) we observed in $[\text{DA}]_{\text{EC}}$. Note that the saturating concentration for DATs is estimated to

![Figure 4](image_url)

**Figure 4.** Results from global sensitivity analysis performed for kinetic parameters representing DAT conformational dynamics. Yellow dots represent the default parameters used in the present study, and blue dots show the results obtained by using as input random combinations of the parameters ($k_{12}$, $k_{23}$, $k_{34}$, $k_{41}$, and $k_{14}$) and their ratios. Results for $k_{34}$, which are very similar to those for $k_{23}$, are not shown. The red curves indicate the mean values and the standard error for successive bins of width 0.5. The Spearman correlation coefficients are –0.71, 0.01, –0.01, –0.44, 0.44, 0.63, and –0.48, in the six respective panels.

![Figure 5](image_url)

**Figure 5.** Time evolution of DA concentration and DAT conformational states averaged over 140 independent MCell runs. **A**, Extracellular DA concentration, $[\text{DA}]_{\text{EC}}$. The average concentration reached under steady-state conditions is 7.8 nM, and the standard deviation of the concentration is indicated by the shaded region is $[\text{DA}]_{\text{EC}} \approx 10$ nM. **B**, The number of DAT molecules, in the unbound inward- or outward-facing state (IF: red; OF: blue), denoted as $N(\text{IF})$ and $N(\text{OF})$, respectively, as a function of time. **C, D**, The average numbers of DATs in substrate-bound OF* and IF* states, $N(\text{OF}^*)$ and $N(\text{IF}^*)$, respectively. The light blue bands show the variance observed in multiple simulations.
Simulations yielded a substrate concentration at half-maximal rate, $K_m$ value, of 2.2 $\mu$M, which falls within the broad range of reported experimental values of 50 nM to 6.6 $\mu$M (Prasad and Amara, 2001; Beuming et al., 2008; Rao et al., 2013). These data confirm that MCell model and simulations reproduce macroscopic quantities consistent with observables such as the average DA concentration in the EC region at half-maximal rate and the overall turnover rate. Next, we make a closer examination of microscopic properties.

**DAT conformers reach a dynamic equilibrium within 100s of milliseconds**

First, we examined the equilibration of the simulated system under uniform surface distribution of DAT molecules on the axonal membrane. The four snapshots in Fig. 6 (and Video 1) illustrate the initial DA release events and the gradual equilibration of the conformational states of DATs. All DAT molecules are assumed to be in the OF state at $t = 0$ (white dots on the surface of the terminals). Simulations start with a first release event (at AZ 1; Fig. 6A), followed by firings with Poisson distribution. The released DA molecules (red dots) rapidly diffuse to the vicinity of the release site, as illustrated in the snapshots at $t = 1$ and 5 ms (Fig. 6B,C). At $t = 700$ ms, we observe a broad spatial distribution of DA (Fig. 6D).

Fig. 6D shows that most of DATs reside in the IF state (colored green) at $t = 700$ ms. This is consistent with the equilibrium probabilities of the four DAT conformers (19.86% OF, 79.90% IF, 0.05% OF+, and 0.19% IF+), which is reached within 500 ms. Fig. 5B–D displays the time evolution of the population of the different states of DAT, averaged over 140 independent runs of 10-s duration each. Most of the DATs fluctuate between unbound OF and IF states, whereas the bound states (OF+ and IF+) are short-lived. Because of their scarcity, the numbers of DATs in IF+ and OF+ states show significant fluctuations during simulations (indicated by the light blue band in Fig. 5C,D).

**DA levels exhibit large fluctuations depending on AZ structure and DAT surface distribution**

We present in Fig. 5A the time course of the DA concentration in the EC region, $[DA]_{EC}$, averaged over 140 runs, which exhibited a standard deviation of $\sim 10$ nM about the mean value of 7.8 nM. These are global fluctuations, i.e., they refer to the average behavior of the
overall microenvironment simulated by MCell. A look at local time- and space-resolved patterns of DA levels, on the other hand, reveals even broader fluctuations depending on the AZ and the particular space or time window examined. The curves in Fig. 7A display the time evolution of the local concentration, $[DA]_{local}$, within a spherical shell of 1.5-$\mu$m thickness, with respective inner and outer radii of 1 and 2.5 $\mu$m (i.e., $1 < r < 2.5 \, \mu m$) centered at the release site, after a release event from the release site on each of the 6 AZs. Results represent the averages over multiple release events (140 per AZ) observed within the time interval $0 < t < 40$ ms.

A considerable variation in the maximum level of DA reached and in the succeeding decay rate is observed in Fig. 7A, depending on the AZ that discharges the DA molecules. For AZ 2 and AZ 3, $[DA]_{local}$ can reach 20–25 nM, but for AZ 6 it remains $<7$ nM. AZ 2 and 3 are on a surface-exposed region of axon terminal 2, in close proximity of non–DAT-expressing cells (Fig. 1C). AZ 6, however, lies inside a cavity on axon terminal 4 that harbors a large density of DATs (Fig. 1C), hence the rapid depletion of the released DA. Other AZs (1, 4, and 5) show an intermediate behavior, because although they are exposed to the EC region, the DAT level in their vicinity is also relatively high (according to fluorescence images).

To examine the effect of neuronal region structural complexity, computations were repeated for a hypothet-

**Video 1.** DA release, diffusion, and reuptake events in the presence at the early (equilibration; 100s of milliseconds) and later (up to 4 s) of MCell simulations. Color code: red, dopamine; green, inward-facing DAT; white, outward-facing DAT. For the first 10 ms, interframe time interval is 1 ms; between 10 and 1000 ms, it is 10 ms; in the rest, 100 ms. To emphasize release events, the frequency of snapshots is reduced between 3.5 and 4 s. Multiple release events are observed, including one at 3.8 s where a closeup view of the corresponding AZ is displayed. [View online]

**Figure 7.** Time evolution of DA levels in the EC region after a single release event in different AZs. A, Time evolution of EC DA levels within a spherical shell of inner radius $r_i = 1 \, \mu m$ and outer radius $r_o = 2.5 \, \mu m$ centered at the AZ or top of the ellipsoid for six different AZs and ellipsoid geometry with same volume and surface area (labeled AZ1–AZ6 and ellipsoid, shown in different colors and patterns; see inset labels). B, C, Local concentration of DA within spherical regions of radius 0.5 $\mu$m (blue) and within spherical shells of $0.5 < r < 1 \, \mu m$ (red), $1 < r < 2.5 \, \mu m$ (green), and $2.5 < r < 5 \, \mu m$ (black), near AZ4 (B) and AZ6 (C), as labeled.
The variations in [DA]_{local} are critically important because, although [DA]_{local} may not be sufficient for activating DA receptors, [DA]_{local} or the corresponding fluctuations Δ[DA] may bring the local DA concentration above the threshold levels required for activating even low-affinity receptors. These results underscore the significance of considering the space-dependent, stochastic nature of DA density fluctuations for estimating the probability of DA receptor activation in complex microenvironments.

**Table 3. Structural properties near AZs**

| Active zone | Void fraction | Tortuosity | Distance to closest DAT (μm) |
|-------------|---------------|------------|-----------------------------|
| 1           | 0.35          | 1.22       | 0.20                        |
| 2           | 0.22          | 1.38       | 0.21                        |
| 3           | 0.20          | 1.56       | 0.24                        |
| 4           | 0.31          | 1.28       | 0.20                        |
| 5           | 0.37          | 1.36       | 0.89                        |
| 6           | 0.36          | 1.23       | 1.22                        |

*Within a spherical region of 2.5-μm radius centered around the release site.

Secondly, the ES region with a simplified (ellipsoidal) geometry that preserved the same EC volume and neuronal surface area as those of the simulated system, together with the same number of DAT molecules on the surface, thus maintaining the same DAT surface density as the above system. DA reuptake evaluated for the same spherical shell (Fig. 7A, dashed black curve) centered around the release site was observed to be faster than that occurring in the realistic environment. The faster removal of DA molecules is due to the more efficient diffusion of DA in this hypothetical environment devoid of irregularities/obstacles.

To make a further quantitative assessment of the role of the AZ geometry and local structural heterogeneity in affecting the different time evolutions of DA clearance observed in Fig. 7A, we examined the tortuosity and void fraction in the vicinity (within 2.5-μm radius) of each of the 6 AZs. The results are presented in Table 3. Both properties affect [DA]_{local}; the clearance of DA is less efficient when the void fraction is low, and the tortuosity is high (e.g., AZ 3 followed by AZ 2). Although void fraction appears to have a dominant effect, tortuosity becomes an important factor when distinguishing the rise time and peak height between AZs that exhibit similar void fractions. AZ 6 has a high void fraction and a low tortuosity, which leads to a rapid reuptake of DA. Another variable we examined is the distance to the closest DAT (Table 3), but it does not have a substantial effect because the diffusion is rapid.

Overall, these results show that (1) the different time evolution of [DA]_{local} in response to DA release from different AZs can be traced back to differences in tortuosity and void fractions and (2) the adoption of uniform tortuosity and void fraction for all AZs would not provide a realistic representation of the heterogeneous synaptic structure, nor would it account for the differential reuptake efficiency originating from these local geometric effects.

Fig. 7B,C further illustrates the variations in [DA]_{local} within gradually increasing distance ranges with respect to the release sites in AZs 4 and 6. [DA]_{local} within a spherical volume of radius r = 0.5 μm remains >10 μM for ~1 ms after DA release from AZ 4, and that within 0.5 ≤ r < 1.0 μm rises more slowly and temporarily reaches 1 μM (Fig. 7B). Comparison with AZ 6 shows the lower [DA]_{local} near the release site (r < 0.5 μm) and the depletion of DA at t ~ 100 ms (Fig. 7C). The large variations in [DA]_{local} due to stochasticity and heterogeneity of the release sites occur during the entire course of the simulations.
Phasic firing favors high transient levels in [DA] while retaining the average [DA]_{EC}

We investigated the effect of distinctive release patterns, tonic and phasic (Goto et al., 2007), on DA reuptake efficiency. Tonic firing was implemented as random spikes generated from a Poisson distribution at 4-Hz frequency (Dreyer et al., 2010), and phasic firing consisted of a burst phase for 0.25 s at 20-Hz frequency followed by a pause of 1 s (Fig. 9A), both yielding the same average frequency. The time evolution of [DA]_{EC} was examined under those two firing patterns for five different models: a well-mixed system (Fig. 9B) and four spatially realistic models derived from image data (cases 1–4 in Fig. 2), which differ in the membrane distribution of DATs. The cases are (1) uniform (Fig. 9C), (2) nonuniform (Fig. 9D), and (3 and 4) sharply nonuniform with two different localizations of high-density regions (Fig. 9E,F; see Table 2 for corresponding surface densities). Each panel displays 140 curves, each corresponding to an independent run.

Tonic and phasic firing patterns elicit markedly different fluctuations in [DA]_{EC}. Tonic firing shows irregular fluctuations (left panels in Fig. 9B–F and Video 2), which usually remain <60 nM in Fig. 9B–D, whereas phasic firing in the same cases (right panels and Video 3) easily exceeds 60 nM, especially when the surface density of DATs is nonuniform (Fig. 9D). Sharply nonuniform distributions of DATs (Fig. 9E,F) give rise to increased [DA]_{EC}, while the difference in the fluctuation behavior of [DA]_{EC} under tonic and phasic firing persists. These results suggest that phasic firing could more readily favor low-affinity DA receptor activation, although the average [DA]_{EC} values over the entire duration of simulations are comparable (Table 4), consistent with the same average firing frequency shared between the five cases.

A closer examination of the transient DA levels within the synaptic region reveals the difference between phasic and tonic firing. We defined [DA]_{syn} as the transient DA level within a sphere of 0.5–μm radius centered at a release site and examined [DA]_{syn} after a single release event. Fig. 10 displays the mean (central dark curve) and the standard deviation (light curves and shade) in [DA]_{syn} for tonic (Fig. 10A,C) and phasic (Fig. 10B,D) firing, under uniform (case 1; Fig. 10A,B) and nonuniform (case 2; Fig. 10C,D) distributions of DATs. Similar to the results for the overall EC (synaptic and extrasynaptic) region (Fig. 9C,D), phasic firing temporarily leads to a higher accumulation of DA in the synapse compared to tonic, which can more readily activate the high-affinity DA receptors (the dashed line indicates the threshold concentration, 10 nM, for their activation).

Finally, we computed the fractional occupancy of low-affinity receptors (Vivo et al., 2006; Casado et al., 2009) during the burst and pause periods of phasic firing and overall period of tonic firing. The results summarized in Table 5 show that the probability of reaching the threshold DA level of 1 μM for binding low-affinity DA receptors is enhanced by a factor of 2–5 in the burst phase of phasic firing, compared with tonic firing. Further computations by adopting a Michaelis–Menten type occupancy model with half-maximal substrate concentration (K_m) of 10 nM for high-affinity receptors and 1 μM for low-affinity receptors confirmed the same behavior. In the latter model, the occupancy of DA receptors as a function of [DA]_{local} is given by

$$\theta([DA]_{local}) = \frac{[DA]_{local}}{K_m + [DA]_{local}}$$

and the results in the last column of the Table 5 are...
obtained on integration over snapshots at 1-μs intervals in 140 trajectories.

Nonuniform surface distribution of DATs is a major modulator of the strength and intensity of DA signaling

Results in Fig. 9 and Table 4 reveal the strong dependence of DA clearance efficiency on DAT surface distribution. DA reuptake is significantly more efficient with a uniform surface distribution of DATs than with a nonuniform distribution, as evidenced by the relative heights of the peaks at low concentrations in the histograms displayed in Fig. 9G,H and the average [DA]EC values listed in Table 4. We investigated the differences among well-mixed, simple spatial (ellipsoid), and complex realistic geometries while keeping the EC volume, surface area, and total density of DAT fixed. Results suggest that the approximation of the axonal morphology by a simplified (ellipsoidal) geometry yields results comparable to those predicted by a well-mixed model. Inclusion of realistic geometry, on the other hand, leads to a broad range of results depending on the specific AZ (Figs. 7 and 8) and the surface distribution of DATs (Fig. 9G,H).

The most efficient clearance is observed in MCell simulations with uniform distribution of DAT. The bimodal distribution of DATs, on the other hand, shows a distinctive distribution of [DA]EC skewed toward higher concentrations, or a significantly suppressed DA reuptake efficiency, as can be seen more clearly in the insets, irrespective of the firing pattern (Fig. 9G,H). This effect becomes more pronounced with increasing heterogeneity of DAT distribution, i.e., going from case 2 to cases 3 and 4.

For a more critical assessment of the duration and intensity of excitatory stimulation induced in response to the two firing patterns, and under different spatial distributions of DATs, we analyzed the peak heights and widths in Fig. 9B–F. The results in Fig. 11 confirm that DAT spatial distribution is a major determinant of dopaminer-
gic signaling strength and duration, whereas the effect of different firing patterns is moderate. In particular, the histograms of the peak widths in Fig. 11A, B, and peak heights in Fig. 11C, D clearly show that bimodal DAT distribution (black and magenta curves) leads to more sustained and stronger excitations compared to uniform (or well-mixed) cases (green and blue curves), originating from less efficient removal of DA. Note that the bimodal distribution was selected to mimic the physiologic density heterogeneities observed in high-resolution images (Fig. 1 and Table 2). The effect of DAT density heterogeneity, manifested by an overall suppression in DA reuptake, is further accentuated in sharply heterogeneous distributions of DATs (cases 3 and 4). The most probable peak heights in those cases are of the order of hundreds of nanomoles per liter and may result in neurotoxicity. Taken together, these results demonstrate that heterogeneity of DAT density can modulate DA signaling.

A comparison between cases 3 and 4 further shows that the placement of DATs in the close neighborhood of the AZs does not increase clearance. The diffusion of DA is fast enough to sample extrasynaptic regions, and the localization of DATs with respect to the AZ has a minimal effect. These results suggest that the mechanism of local clearance as envisioned by the gatekeeper model (Rothstein et al., 1994) does not necessarily increase reuptake efficiency.

To examine the change in DA reuptake behavior possibly induced by DAT 2D displacement, we repeated the simulations by allowing DAT 2D diffusion with a coefficient of $3 \times 10^{-10}$ cm$^2$/s on the plasma membrane (Fig. 12). Comparison of the initial and final (at the end of 10-s simulations) averaged over 140 independent runs showed that no significant change occurred in the position of

![Video 2. Dynamics of DA molecules and time evolution of $[DA]_{EC}$ under tonic firing. Same color code as in Video 1. The total duration of the movie is 5 s. The bottom plot displays the time evolution of $[DA]_{EC}$ during the course of simulations. [View online]](image)

![Video 3. Dynamics of DA molecules and time evolution of $[DA]_{EC}$ under phasic firing. Same color code as in Video 1. The time span of the movie is 5 s. [View online]](image)

| Space DAT distribution | $[DA]_{EC}$ (nM) | $\Delta [DA]_{EC}$ (nM) |
|------------------------|-----------------|-------------------------|
| Tonic                  | Phasic          |
| Well-mixed DA$^a$      | 8.16 ± 0.92     | 8.29 ± 1.41             |
| Ellipsoid geometry     | 8.21 ± 0.85     | 8.33 ± 1.44             |
| Complex space          |                 |                         |
| Case 1 Uniform$^b$     | 7.54 ± 0.86     | 6.93 ± 1.92             |
| Case 2 Nonuniform$^c$  | 16.59 ± 1.26    | 14.19 ± 2.44            |
| Case 3 Sharp bimodal$^d$| 33.12 ± 3.13    | 29.65 ± 4.49            |
| Case 4 Sharp bimodal$^d$| 32.49 ± 2.86    | 32.89 ± 3.68            |

$^a$Space/structure-independent. All other cases are in realistic DA neuronal geometry.

$^b$DAT densities for uniform and nonuniform (bimodal) distributions; see Table 2.

$^c$Case 2 ($p_h = 6339/\mu m^2$ and $p_l = 50/\mu m^2$).

$^d$Cases 3 and 4 ($p_h = 30,000/\mu m^2$ and $p_l = 50/\mu m^2$; see Figs. 2 and 11).
DATs, owing to the slow diffusion of DATs compared with the time scale of simulations. Overall, these results show that increasing heterogeneity in the surface distribution of DAT leads to lower DA clearance efficiency and/or stronger and more sustained signaling.

**Discussion**

**Overview**

In the present study, we reconstructed *in silico* the 3D structure of DA terminals, the location of AZs, and the

**Table 5. Percentage of high- and low-affinity DA receptors that can potentially bind DAs**

| DAT distribution | DA receptor type | Firing pattern and stage | Percent probability of DA binding to DA receptors | Percent probability of DA binding to DA receptors |
|------------------|------------------|--------------------------|-----------------------------------------------|-----------------------------------------------|
| Uniform (case 1) | High affinity    | Tonic                    | 40.26                                         | 45.98                                         |
|                  |                  | Phasic burst             | 45.09                                         | 51.80                                         |
|                  |                  | Phasic pause             | 4.54                                          | 50.07                                         |
|                  | Low affinity     | Tonic                    | 0.40                                          | 2.22                                          |
|                  |                  | Phasic burst             | 1.10                                          | 4.64                                          |
|                  |                  | Phasic pause             | 0.06                                          | 1.92                                          |
| Nonuniform (case 2) | High affinity | Tonic                    | 71.14                                         | 63.84                                         |
|                  |                  | Phasic burst             | 65.88                                         | 62.70                                         |
|                  |                  | Phasic pause             | 72.05                                         | 64.5                                          |
|                  | Low affinity     | Tonic                    | 0.35                                          | 3.47                                          |
|                  |                  | Phasic burst             | 1.97                                          | 5.47                                          |
|                  |                  | Phasic pause             | 0.03                                          | 2.97                                          |

*The burst period is 2.5–2.75 s after DA release, and the pause 2.75–3.75 s (see Fig. 10).*

Figure 10. Time evolution of DA level in the synapse after a release event. **A–D**, [DA]syn as a function of time for tonic (left) and phasic (right) firing, under uniform (top) and nonuniform (bottom) distribution of DATs. The mean and variation in [DA]syn within 1 μm collected from 140 independent runs are displayed. Results are shown for the time interval 2.5–3.75 s as a single firing block, representative of dynamic equilibrium reached under stationary conditions. The thick line in the middle of cloud is the mean value over all runs, and the cloud represents the standard deviation, highlighting the large fluctuations in the synaptic DA levels.
spatial distribution of DATs based on electron and fluorescence microscopy data with transgenic mice (Block et al., 2015) and conducted a series of MCell simulations under different conditions. We determined the spatiotemporal distribution of DA molecules in the synapse and extrasynaptic regions in response to different firing pat-
terns and in the presence of different surface distributions of DATs. These simulations show the effects of the synaptic and extrasynaptic morphology, as well as the heterogeneity of DAT surface distribution (as observed in experiments), on the overall efficiency of DA clearance and the probability of activating high- and low-affinity receptors. Heterogeneous distribution of DATs is shown to reduce the clearance efficiency, the effect being sharper with increased heterogeneity, and not affected by DAT 2D diffusion on the plasma membrane. Although DA dynamics is diffusion-controlled in general, a global sensitivity analysis indicated that the DA-binding rate of DAT and the relative population of its outward- and inward-facing conformers in the apo state (denoted as OF and IF) had major effects on DA reuptake efficiency. Overall, the results underscore the utility of conducting spatiotemporally realistic simulations in elucidating the dependence of dopaminergic signaling on both the surface distribution and conformational mechanics of DATs.

Recent MCell simulations of glutamatergic signaling based on EM images of hippocampal neuropil for investigation of the effect of spatial heterogeneities (Kinney et al., 2013; Bartol et al., 2015a, c) also indicated the importance of including the microenvironment for an accurate description of the electrophysiology and biochemistry of neurotransmission events. No such simulations had been performed for DA neurons to date, mainly because of the lack of high-resolution image data of DA terminals for reconstituting in silico the simulation environment. Recent advances in imaging DA neurons (Block et al., 2015) permitted us to overcome this barrier. Furthermore, improved understanding of the structural dynamics of DAT (Cheng and Bahar, 2015; Cheng et al., 2015; Gur et al., 2017) helped us build a simplified kinetic scheme (Fig. 3) that we adopted in MCell simulations.

**DA reuptake simulations require spatially extended (~10^-9-µm^2) structural models**

The simulation of dopaminergic signaling events presents new challenges compared with that of neurotransmission in glutamatergic synapses. The transporter cycle rate in the latter case is ~35 glutamate molecules/s per glutamate transporter, whereas the rate in dopaminergic neurons is 1–5 DA molecules/s per DAT (Wadiche et al., 1995; Povlock and Schenk, 1997; Prasad and Amara, 2001; Rice et al., 2011). Thus, simulation of DA clearance requires longer computing times. In addition, glutamate transporters are localized in pre- and postsynaptic regions and mostly in the surrounding glia in the close neighborhood of the AZ (Seal and Amara, 1999; Danbolt, 2001). DATs, on the other hand, are preferentially located in the extrasynaptic regions of the axons and dendrites, and most of the reuptake events take place at sites distal from the AZs (Nirenberg et al., 1996; Rice et al., 2011; Block et al., 2015). Realistic simulations of such spatially distributed events require the adoption of model systems composed of multiple synapses, multiple AZs, and heterogeneous distributions of multiple DAT clusters. We reconstructed in silico a relatively large (10 × 10 × 7.2-µm) striatal region.

**Spatial irregularities and hindrance in the interstitial region between neurons limits DA receptor activation**

Our simulations show that under conditions that reproduce physiologic levels of DA and turnover rates (Prasad and Amara, 2001; Beuming et al., 2008; Rao et al., 2013), the complex geometry of DA terminals modulates the occupancy of high-affinity DA receptors to <100%, even under sustained, elevated stimulation conditions (Fig. 8A–C), consistent with the concept of a dead space (Kamali-Zare and Nicholson, 2013). In contrast, less detailed partial differential equation models (Dreyer et al., 2010) and algebraic models (Cragg and Rice, 2004) predict full occupancy of high-affinity receptors under the same conditions. This difference invites attention to the effect of EC spatial irregularities on dopaminergic signaling efficiency in the brain (Syková and Nicholson, 2008).

**Heterogeneous surface distribution of DAT reduces the effectiveness of DA clearance**

EM and fluorescence images showed that the distribution of DAT is heterogeneous in different parts of the striatum (Block et al., 2015). Recent studies indicate that the population of DATs may vary depending on the membrane curvature (Caltagarone et al., 2015). Notably, selected DAT mutants that have disrupted OF conformations do not accumulate in filopodia, suggesting that access to the OF/OF* state may be a prerequisite for DAT to populate the filopodia (Caltagarone et al., 2015). Further examination showed that binding of cocaine and its fluorescent analog JHC1-64 also alters the plasma membrane distribution of both wild-type and mutant DATs. Cocaine binding arrests DAT in the OF state, and cocaine-bound DATs predominantly localize in the filopodia (Ma et al., 2017). Likewise, zinc binding, also known to stabilize the OF state, led to an increase in the level of DAT mutants in the filopodia. These observations suggest that the membrane curvature is a determinant of DAT clustering, with the convex shape of the filopodia favoring the localization of OF DATs. Furthermore, clustering of OF DATs could remodel the membrane to induce an overall outward bending (Ma et al., 2017). The present study suggests that such targeting of axonal protrusions by OF DATs, along with their membrane-remodeling capacities, may regulate DA reuptake.

The total number of DAT molecules was sufficiently large in current simulations (~200,000) to ensure normal DA clearance consistent with physiologic levels. However, simulations repeated with the same number of DATs distributed nonuniformly on the neuronal membranes showed a reduction in reuptake capacity that became more pronounced with increasing surface density heterogeneity. The surface distribution of DATs indeed emerged as a major determinant of the efficiency of reuptake, or conversely, the strength and duration of excitatory signaling (Figs. 9 and 11).

In principle, one might expect that the increased population of DATs in dense regions would counterbalance the effect of lower surface area coverage on DA reuptake efficiency. However, DA diffuses fast enough to escape from these dense regions after a few encounters, resulting
in reduced DA reuptake. These observations further support the importance of adopting a realistic representation of the heterogeneity of DAT surface distribution, especially with low copy numbers of neurotransmitters, for a realistic assessment of DA reuptake capacity.

**Firing patterns determine the relative levels of inhibitory versus excitatory responses**

Our simulations revealed that the average [DA]_{EC} exhibited little dependence on firing pattern, provided that the average firing frequency was maintained, but the local levels within a synapse or close to DA release sites exhibited strong dependencies. The size of the fluctuations, Δ[DA], in DA levels were relatively small in tonic firing but large in phasic firing, especially at the burst phase and under sustained phasic firing (Fig. 9B–F). Low-affinity DA receptors need two orders of magnitude larger DA concentrations than high-affinity receptors to get activated. As a result, the activation of low-affinity receptors is limited, if not unlikely, with tonic firing (Table 5). Phasic firing, on the contrary, induces a larger variance in both global and local DA concentrations and is two to five times more likely to activate low-affinity receptors during the burst phase of DA signaling (Table 5). Previous work showed that the D1 receptors are mostly localized at the postsynaptic membrane (Cadet et al., 2010), and our simulations show that only those low-affinity receptors localized near the synapse can be activated. In the case of high-affinity receptors, on the other hand, proximity to release site is not a requirement for being activated: high-affinity receptors located on distal regions may be equally activated.

DA receptors regulate the activity of DA neurons. One distinction between high- and low-affinity DA receptors is the type of response they elicit in the cell, inhibitory or excitatory (Keeler et al., 2014), after their activation on DA binding. D1-like receptors, which are usually low-affinity receptors, are involved in excitatory signaling, whereas D2-like receptors trigger inhibitory signaling processes. High-affinity receptors may be activated by both phasic and tonic firing. However, our simulations suggest that low-affinity receptors would be rather activated on phasic firing.

Among D2-like receptors, D2 autoreceptors are known to be key regulators of DA transmission, located on most axon terminals (Ford, 2014). Although D2-like receptors are usually high-affinity receptors, D2 autoreceptors exist in both high- and low-affinity states, and recent studies indicate that the functionally relevant D2-autoreceptors are predominantly in low-affinity state. Their inhibitory action is particularly important to suppressing DA release from presynaptic cells under prolonged bursts of action potentials. They suppress DA release in the striatum by several mechanisms, e.g., by inhibiting the voltage-gated calcium channels that trigger exocytotic DA release or increasing DAT activity or surface expression (Ford, 2014). Previous studies suggested that tonic firing does not raise [DA]_{EC} to levels sufficiently high to activate D2-autoreceptors (Beckstead et al., 2007; Ford, 2014). The low probability of binding low-affinity receptors under tonic firing demonstrated in the present simulations is consistent with the functioning of D2-autoreceptors as low-affinity receptors.

A previous model modified the probability of DA release on an action potential according to high-affinity receptor occupancy (Dreyer and Hounsgaard, 2013). In our current model, receptor occupancy has not been coupled to DA release and reuptake. However, our modeling framework allows for implementing a mechanistic model that incorporates the mediation of DA release and DAT expression by D2-autoreceptors. Such an extension may be key to improved interpretation of DA reuptake dynamics at longer time scales and under disease states.

**The modeling framework is extensible to analyzing the effect of psychostimulants on DA reuptake dynamics**

DAT kinetics is often described by the alternating access model (Vaughan and Foster, 2013). Most cell-level models of DA transport at the cellular level assume an immediate transition from IF to OF state right after DA release to the cell interior (Cragg and Rice, 2004; Rice and Cragg, 2008; Sulzer et al., 2016), and the transition from OF to IF in the unbound form is not explicitly considered. In contrast, molecular simulations revealed a complex dynamics (Cheng and Bahar, 2015), which we reduced to a four-state kinetic model (OF, IF, OF*, and IF†). The translocation and release events after substrate binding are much faster compared with the return of apo IF state to OF state and succeeding substrate binding event, which depends on prior probabilities of encounters between DAT and DA molecules. These features result in short lifetimes for the bound forms of DAT and the dominance of the relative populations of the unbound OF and IF states in determining the overall clearance rate (Fig. 5). In particular, the balance between the OF and IF states (or the ratio of the associated rate constants, k_{14}/k_{41}, using detailed balance principle) emerged as a major determinant of reuptake efficiency, as it directly defines the fraction of reuptake-ready (OF) DATs.

Previous x-ray crystallographic studies (Penmatsa et al., 2015; Wang et al., 2015) as well as structure-based simulations (Cheng et al., 2015) have shown how the conformational state of DAT and substrate binding affinity may change in the presence of different antidepressant and psychostimulants drugs, such as cocaine and amphetamine (AMPH). Antidepressants usually arrest DAT in OF state [and the same effect has been observed for the homolog serotonin transporter in the presence of antidepressants (S)-citalopram or paroxetine (Coleman et al., 2016)], thus preventing the progress of the transport cycle, and causing an increase in EC DA levels. On the other hand, the occupancy of the Zn^{2+}-binding site (Stockner et al., 2013; by zinc or other transition metals) reduces the DA binding affinity of DAT (Li et al., 2017). The current modeling framework can readily be extended to incorporate such effects through suitable readjustment of kinetic parameters in Fig. 3.

AMPH, on the other hand, appears to be a substrate for DAT that competes with DA for transport (Zhu and Reith, 2008). AMPH is also known to promote both DA efflux...
(reverse transport from the cell interior to the EC region) and DAT internalization (Wheeler et al., 2015). The current simulations focused on the influx of DA from the EC region to the presynaptic cell interior, but the modeling framework is readily extensible to simulating DA efflux as well. Other extensions of the present framework could incorporate the effects of DAT internalization, membrane polarization (Richardson et al., 2016), or perturbations at the membrane-proximal N-terminal residues (Sorkina et al., 2009). Thus, the present study opens the way to quantitative modeling of the effects of antidepressants, psycho-stimulants, and substances of abuse on the deregulation of DA signaling.

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