

## Molecular Dynamics Study of Non-nucleoside Reverse Transcriptase Inhibitor 4-[[4-[[4-(*E*)-2-Cyanoethenyl]-2,6-dimethylphenyl]amino]-2-pyrimidinyl]amino]benzotrile (TMC278/Rilpivirine) Aggregates: Correlation between Amphiphilic Properties of the Drug and Oral Bioavailability

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The non-nucleoside reverse transcriptase inhibitor (NNRTI) TMC278/rilpivirine is an anti-AIDS therapeutic agent with high oral bioavailability despite its high hydrophobicity. Previous studies established a correlation between ability of the drug molecule to form stable, homogeneous populations of spherical nanoparticles (~100–120 nm in diameter) at low pH in surfactant-independent fashion and good oral bioavailability. Here, we hypothesize that the drug is able to assume surfactant-like properties under physiologically relevant conditions, thus facilitating formation of nanostructures in the absence of other surfactants. The results of all-atom molecular dynamics simulations indeed show that protonated drug molecules behave as surfactants at the water/aggregate interface while neutral drug molecules assist aggregate packing via conformational variability. Our simulation results suggest that amphiphilic behavior at low pH and intrinsic flexibility influence drug aggregation and are believed to play critical roles in the favorable oral bioavailability of hydrophobic drugs.

Good oral bioavailability is increasingly considered to be a key feature of new chemical entities (NCEs<sup>d</sup>) as potential drug candidates identified through high throughput screening (HTS), combinatorial chemistry, and cell-based activity assays.<sup>1</sup> Current statistics report that 35–40% of NCEs are known to belong to the biopharmaceutics classification systems (BCS) class II type of molecules with poor solubility and high permeability properties.<sup>1,2</sup> Attempts to improve oral bioavailability for class II compounds focused on enhancement of drug solubility via a myriad of formulation designs including salt complexation,<sup>3</sup> prodrug formation,<sup>4</sup> particle size reduction,<sup>5,6</sup> self-emulsifying drug delivery system (SEDDS),<sup>6–9</sup> self-microemulsifying drug delivery system (SMEDDS),<sup>6,9</sup> micellar/surfactant system,<sup>10</sup> and solid dispersion.<sup>1,11,12</sup> Most of these formulation designs focus on generation of stable, nanosize delivery units obtained either by altering the physicochemical properties of the drug molecules or by addition of cosolvents, surfactants, and/or lipid suspensions. Unfortunately, besides the intrinsic complexity and

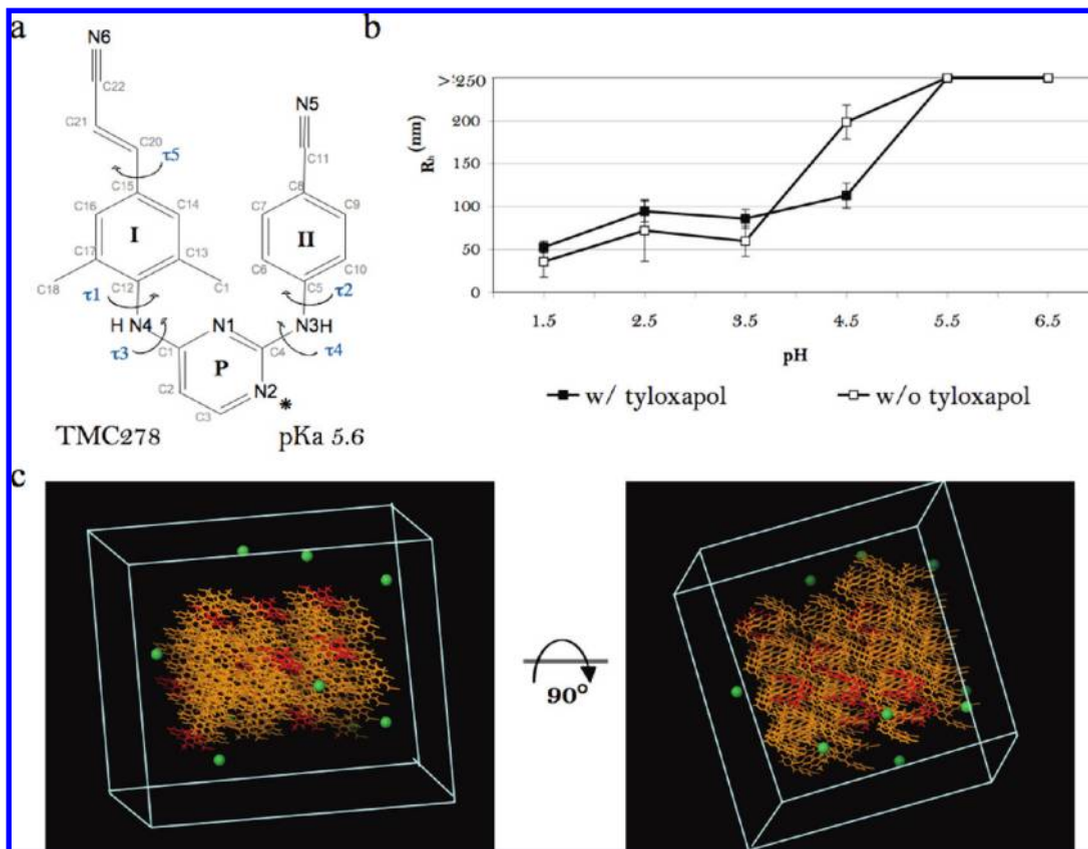
drug specificity involved in implementation of some of the mentioned methods, there are also issues of drug precipitation, low drug loading, instability, excipient-related toxicity, and first-pass metabolism that require further investigation and improvement.<sup>1,13</sup>

Though uncommon, there are NCEs that fall outside of the poor solubility/poor bioavailability trend<sup>14</sup> such as the highly hydrophobic NNRTI TMC278, compound **1** (Figure 1a), which has been shown to be highly potent against HIV-1 at relatively low oral doses (~25–75 mg/day).<sup>15,16</sup> Compound **1** belongs to the diarylpyrimidine (DAPY) class of anti-AIDS drugs that specifically bind to a highly hydrophobic pocket in the reverse transcriptase (RT) enzyme of HIV-1.<sup>15,17,18</sup> Crystallographic and computational studies have shown that ability of the molecules in this class to tolerate prevalent mutations in the binding pocket stem from their capacity to observe multiple binding modes via torsional flexibility (wiggling) and local repositioning (jiggling) while adapting to movements of surrounding pocket residues.<sup>15,19,20</sup> For some of the NNRTIs, exceptional RT-binding properties are accompanied by surprisingly good oral bioavailability.<sup>16,21</sup> While some of the compounds, such as the recently approved DAPY NNRTI drug TMC125/etravirine, require formulation designs involving solid dispersion methodology to improve its intrinsically low oral bioavailability,<sup>22</sup> others, including **1**, were shown to have good bioavailability in animal models and humans with minimal formulation designs.<sup>16,21,23</sup>

On the basis of observations in a standard cell culture assay and biophysical measurements in solutions simulating fasting gastric conditions, it has been proposed that some NNRTIs can form homogeneous populations of small (~100–120 nm diameter) drug aggregates that contain ~500000–2000000

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<sup>d</sup> Abbreviations: NNRTI, non-nucleoside reverse transcriptase inhibitor; NCE, new chemical entity; HTS, high throughput screening; BCS, biopharmaceutics classification system; SEDDS, self-emulsifying drug delivery system; SMEDDS, self-microemulsifying drug delivery system; RT, reverse transcriptase; DAPY, diarylpyrimidine; MD, molecular dynamics; DLS, dynamic light scattering; DPD, dissipative particle dynamics; SPC, simple point charge;  $N_{\text{agg}}$ , number of molecules in an aggregate;  $S_m$ , molecule based solvation parameter; SASA, solvent-accessible surface area; mSASA, maximum solvent-accessible surface area.



**Figure 1.** Surfactant-dependent drug aggregation in vitro and the 10% protonated aggregate model in silico. (a) Molecular structure of **1** with wing I, wing II, and pyrimidine rings labeled as **I**, **II**, and **P**, respectively. (\*) denotes the preferred protonation site at the N(2) atom of the pyrimidine ring, as suggested by the small-molecule crystal structure of protonated compound **1**.<sup>45</sup> (b) Hydrodynamic radius distribution as a function of solution pH for 0.1 mM **1** in the presence and the absence of 0.1% tyloxapol. (c) The starting conformation of the 10% protonated model in a 60 Å × 60 Å × 60 Å cell. Color assignment: yellow, neutral compound **1**; red, charged compound **1**; green, chloride ion. SPC water molecules are not shown.

molecules per particle.<sup>21,23</sup> Oral bioavailability properties of the DAPY compounds were correlated with their lipophilicity, transepithelial transport, human exposure, and ability to form small drug aggregates at low pH and high drug concentration.<sup>23,21</sup> Consequently, it was hypothesized that small drug aggregates were the biological units of delivery that formed at high drug concentrations in the stomach and were being further trafficked into lymphatic circulation by small intestine enterocytes via either paracellular or transcellular routes.<sup>23</sup>

Among currently known nanoparticulate structure types, we have reasons to believe that microemulsion is the most appropriate model to describe NNRTI drug aggregates. Microemulsions are isotropic, thermodynamically stable ternary or pseudoternary systems with 80–100 nm diameter particles, otherwise known as swollen micelles that consist of oil (hydrophobic phase), water (hydrophilic phase), surfactant (stabilizing agent), and frequently a cosolvent.<sup>24–27</sup> Surfactants are known to lower the surface tension between two immiscible phases<sup>24–27</sup> and to have specific structural characteristics such as charged or polar head groups and hydrophobic tails. In the case of some DAPY NNRTIs, we hypothesize that under acidic conditions, protonated drug molecules have surfactant-like features with the protonated pyrimidine ring being a headgroup and the diaryl wings composing the hydrophobic tail. We propose that those NNRTI DAPY compounds that can exhibit surfactant-like properties upon protonation can facilitate microemulsion-like aggregate formation.

Here, we examine compound **1** behavior at the water/aggregate interface using all-atom molecular dynamics (MD) simulations. A number of computational studies have been targeted at water/vapor interfaces<sup>28</sup> and microemulsions such as simulations of single or multiple surfactant molecules in explicit solvent,<sup>29</sup> coarse-grained simulations using self-consistent field theory,<sup>30</sup> and dissipative particle dynamics (DPD) based on nuclei formation and growth via coalescence exchange or coagulation of water-in-oil drops for simulation of microemulsions, micelles, etc.<sup>31,32</sup> However, all-atom simulations of aggregate systems consisting of a half million molecules or more are not currently feasible. Therefore, we have devised a computational approach to study the surface dynamics of a molecule-sized aggregate rather than the full nanometer-sized aggregates. In the present study, we monitor pH-dependent aggregate surface formation with the goal to evaluate the behavior of protonated and neutral drug molecules in terms of their potential surfactant-like behavior.

## Results

**Solution Studies.** On the basis of the earlier reported solution study of pH-dependent aggregation behavior for compounds with good oral bioavailability,<sup>23</sup> we hypothesized that DAPY NNRTIs exhibit surface-active properties under conditions of low pH and therefore are able to form aggregates in a surfactant-independent fashion. Previous results indicated that **1** forms spherical nanostructures with

**Table 1.** Composition and Analysis of the Results of the Computational Models of Compound **1** Aggregates

	0%	10%	30%	60%	90%
computational models					
total no. of drug molecules	100	100	100	100	100
no. of protonated molecules	0	10	30	60	90
no. of Cl <sup>-</sup> atoms	0	10	30	60	90
no. of water molecules	5135	4783	4720	4582	4324
final aggregate composition <sup>a</sup>					
$N_{\text{agg}}$	98	98	85	46	0 <sup>b</sup>
no. of surface molecules	48	48	49	36	0 <sup>b</sup>
no. of core molecules	50	50	36	10	0 <sup>b</sup>
no. of protonated surface molecules	0	9	17	3	0 <sup>b</sup>
SASA (Å <sup>2</sup> )	10610	11663	10728	6950	0 <sup>b</sup>
surface charge density ( $e/\text{nm}^2$ )		0.008	0.016	0.004	0 <sup>b</sup>
U conformation populations <sup>c</sup>					
aggregate core	99.8	65.3	61.8		
aggregate surface, neutral	90.9	60.3	45.6		
aggregate surface, protonated		63.5	58.9		
$L_{\tau 3}$ conformation populations <sup>c</sup>					
aggregate core	0.2	27.9	37.2		
aggregate surface, neutral	9.1	35.9	51.1		
aggregate surface, protonated		28.3	11.2		

<sup>a</sup> Aggregate composition was determined using the solvation parameter  $S_m$  (see Materials and Methods for details). <sup>b</sup> Not applicable due to lack of aggregation. <sup>c</sup> Fraction in percent.

average radii  $\sim 60$  nm in aqueous solution in the presence and in the absence of a surface-active agent at pH 1.5.

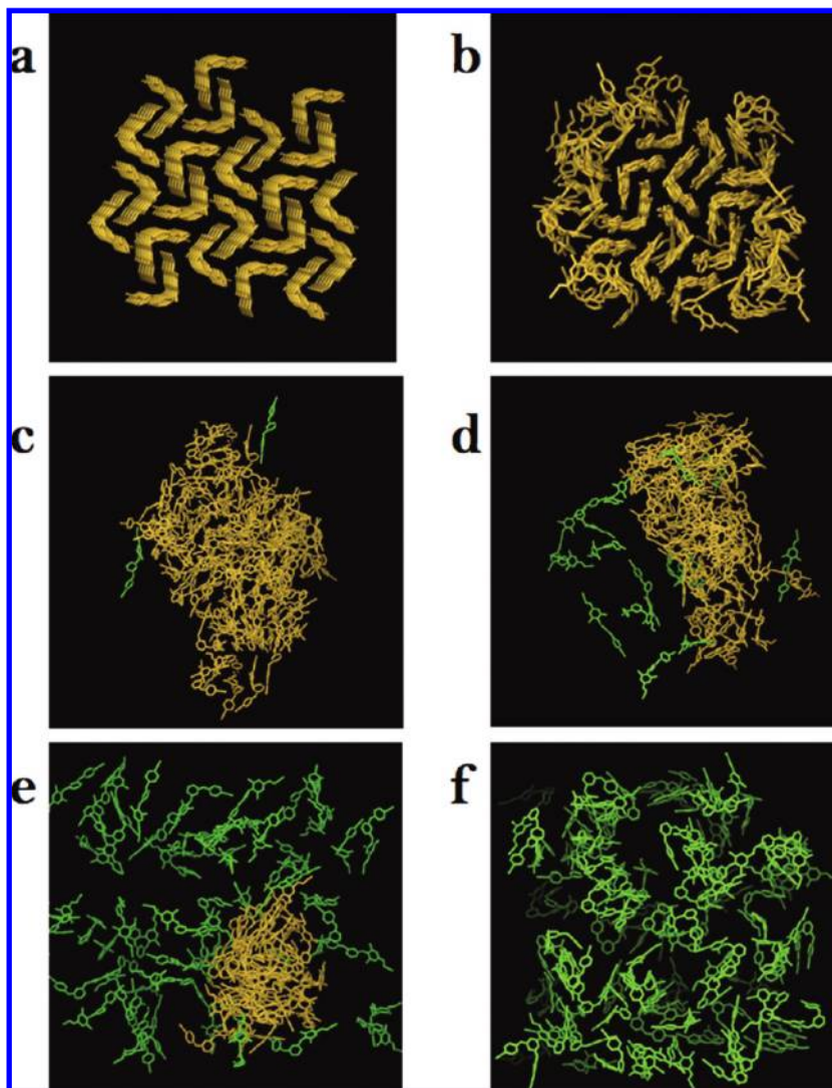
To answer the question of whether or not the aggregation tendencies of the drug depend on the presence of a stabilizing agent at higher pH conditions, we conducted a series of DLS experiments in the presence and absence of the surfactant. These studies show that **1** is able to display surface activity only at low pH. We monitored the aggregation of 0.1 mM drug solutions in the presence and absence of the surfactant tyloxapol (Figure 1b). As shown in Figure 1b, the average size of compound **1** aggregates in aqueous solutions tends to increase with increase in pH. At the most acidic conditions (pH 1.5), slightly larger aggregates were observed in the presence of tyloxapol (average drug aggregate radius  $52.5 \pm 6.8$  nm) than in its absence ( $35.5 \pm 17.7$  nm). This trend consistently continues in the pH range from 2.5 to 3.5, where in the presence of tyloxapol the average aggregate size was greater by  $\sim 20$  nm than in the tyloxapol-free solutions. At pH 4.5, the trend observed at lower pH was reversed and smaller aggregate size was observed for samples with tyloxapol ( $112.5 \pm 14.5$  nm) versus without ( $198.5 \pm 20.0$  nm). Formation of the homogeneous aggregate population at higher pH values was not observed and is best characterized as a precipitate.

On the basis of these observations, we conclude that compound **1** aggregation does not depend on the presence of added surfactant only at low pH conditions. On the basis of the compound **1**  $pK_a$  of 5.6 and observed in vitro aggregation tendencies, we hypothesize that the surface-active agent responsible for the formation of the aggregates in the absence of tyloxapol is the protonated (positively charged) form of **1**. The tendency toward larger aggregates with increasing pH is thus rationalized in terms of the decrease of the fraction of surface active protonated **1** molecules. To understand further the effect of protonation on the drug aggregate assembly processes in aqueous environments, we conducted a series of MD simulations to further probe the protonation-dependent behavior of the drug.

**Computational Studies of Drug Aggregates.** To validate the hypothesis stated above and to unambiguously investigate the pH-dependent aggregation properties of **1**, we have formulated minimalist models of solutions containing only water, compound **1** molecules (of which 0 to 90% were charged), and sufficient chloride ions to neutralize the system (Table 1, Figure 1c). By design, the drug concentration in aggregate models exceeds that of the in vitro conditions yet is believed to be reasonable for studies of aggregate morphology, aggregate packing, conformational variability of drug molecules, and their surface presentation. Morphology, charge distribution, and packing analysis suggest that drug aggregate structure is pH dependent and biphasic with a partially neutral core and a more disordered and partially charged surface.

Visual inspection of the aggregate morphology during the MD simulations of the compound **1**/water systems at the increasing level of protonation indicates that aggregate formation is a pH-dependent phenomenon. The original cubic shape of the starting microcrystal (Figure 2a) is lost for all but the model without protonated **1** molecules (0% protonation). The models with 10 and 30% protonated compound **1** molecules produce a continuous aggregate phase, while those with 60 and 90% protonation result in either partially or fully dissociated solutions. As reported in Figure 2 and Table 1, the 0, 10, and 30% protonated models yield structures with similar  $N_{\text{agg}}$  and SASA, whereas the 60% and 90% protonated models produced structures with significantly smaller  $N_{\text{agg}}$  and SASA values and therefore were not used in quantitative comparison analysis.

Molecular composition analysis of the aggregates reported in Table 1 indicates that charge saturation of the aggregate surface is reached at under 30% protonation. While the number of drug molecules making up the surface of the aggregates in the 10 and 30% protonated models is similar (48 and 49 molecules, respectively), the corresponding number of charged molecules at the surface are different (9 and 17, respectively). Consequently we observe that a

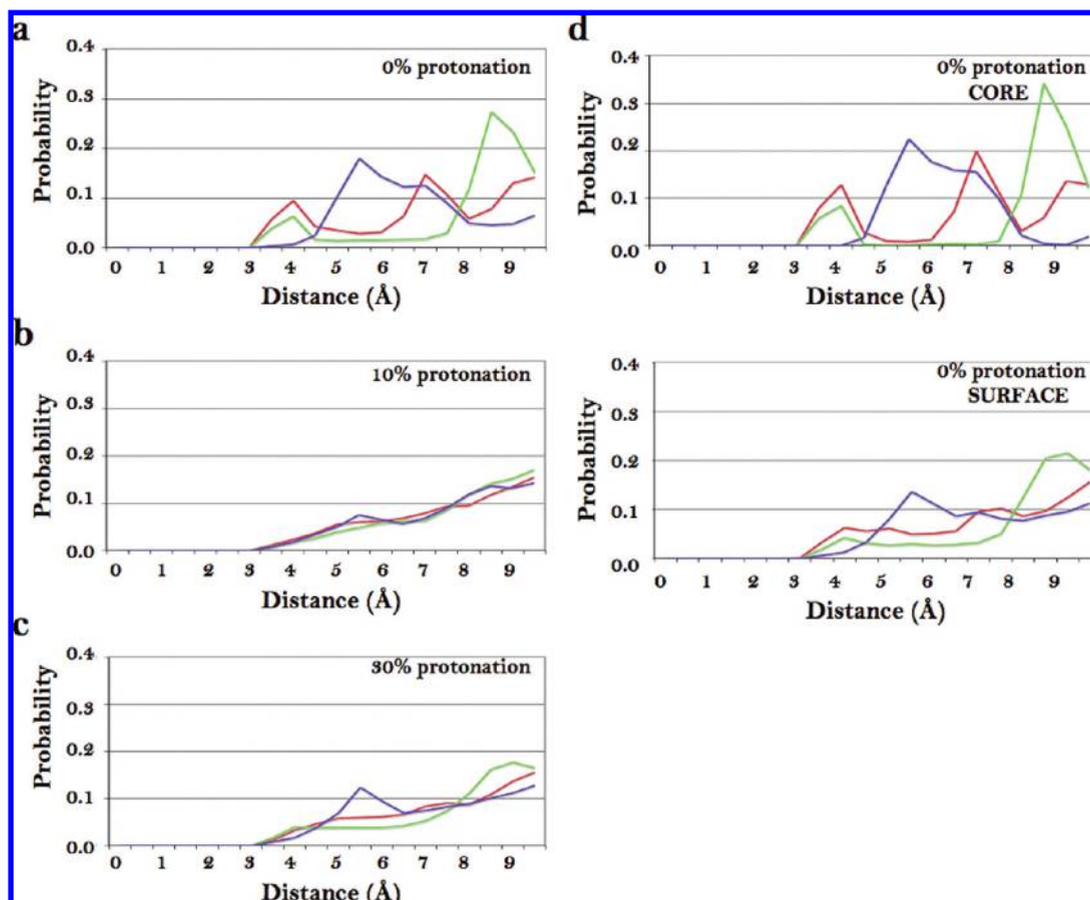


**Figure 2.** Snapshots of compound **1** aggregate structures extracted from the MD trajectories at various levels of protonation. (a) Starting conformation of the 0% protonation model. (b–f) Final conformations of the 0, 10, 30, 60, and 90% protonation models. Water molecules and counterions are not shown. Color assignment: yellow, aggregated compound **1** molecules; green, dissociated compound **1** molecules (aggregate association is measured based on  $S_m$  parameter described in Materials and Methods).

3-fold increase in number of charged molecules (from 10 to 30%) is accompanied by only a 2-fold increase in the surface charge density, from 0.008 to 0.016 charge/nm<sup>2</sup>, respectively. The nonlinear trend in surface charge distribution suggests that system reaches surface charge saturation at under 30% protonation. Furthermore, full dissolution of the aggregate at 90% protonation (Figure 2f) after only 2 ns of MD simulation suggests that under sufficiently acidic conditions aggregate formation would be inhibited.

Visual examination and quantitative packing analysis of the structures obtained from the simulations indicate that the aggregates tend to assume biphasic structures. The structures of the aggregates obtained at 0% protonation represent extreme examples of this phenomenon. In Figure 2b, we see that the aggregate displays two clearly distinguishable layers, one of which has retained the starting crystalline packing and the other, at the water interface, is partially disordered. Interestingly, such clear differentiation is present only at 0% protonation. Even at only 10% protonation (Figure 2c), the core of the aggregate does not display crystalline ordering.

Our results suggest that protonated **1** molecules tend to induce a liquid-like disorder at the water/aggregate interface that in turn induces disorder in the core of the aggregates. The water/aggregate interface of the 10% and 30% protonated models is populated preferentially by protonated molecules and displays greater disorder than the interface of the 0% model. The natural implication of this observation is that crystalline-like ordering would be more prevalent in regions deep in the aggregate core distant from the surface layer if a larger number of drug molecules would be employed in atomistic simulations. This hypothesis is strengthened by the resiliency displayed by the crystalline-like ordered phase at 0% protonation, which, unlike at higher level of protonation, remained stable for the entire duration of the simulation. In conclusion, although the modeling design employed here is not intended for studying the core structure of the aggregates, the present *in silico* results indicate that the nanometer-sized aggregates of **1** we observed *in vitro* likely contain cores largely unaffected by the specific solution pH conditions, which are instead predicted to have a marked effect on the structure of the water/aggregate interface.



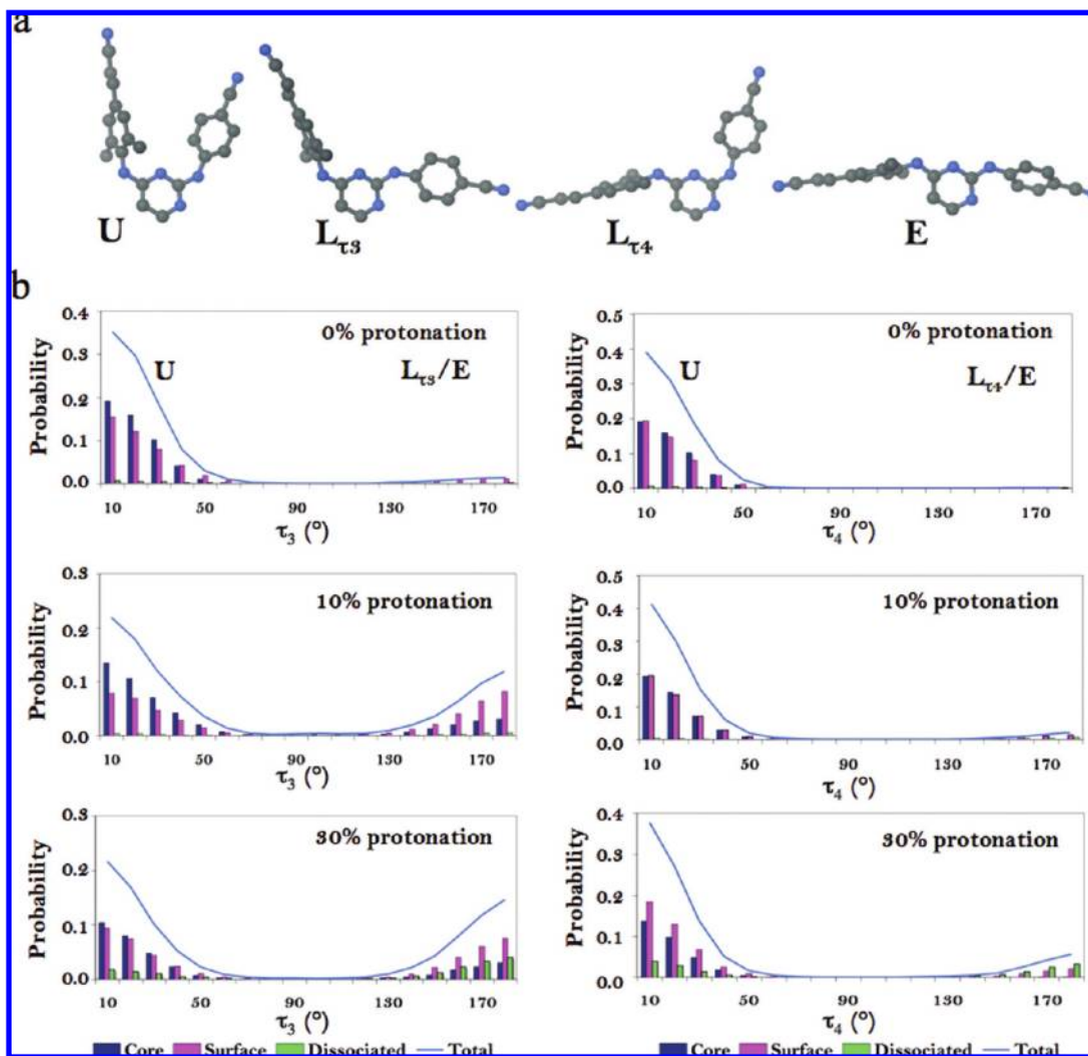
**Figure 3.** Intermolecular ring-to-ring distance distributions for the 0, 10, and 30% protonation models. (a) 0%; (b) 10%; (c) 30%. (d) Ring-to-ring distance distribution functions for the core compound **1** molecules and surface compound **1** molecules from the 0% protonation model. Color assignment: red, distances between ring I and ring II; green, distances between ring II and ring II; blue, distances between ring II and pyrimidine ring.

Ring-to-ring distance distribution analysis of the MD trajectories confirms that compound **1** molecules form a liquid-like layer at the water/aggregate interface, while the core molecules of the 0% model preserve the crystalline arrangement. Distance analysis shows that stacking interactions characteristic of crystal packing are primarily present only in the 0% protonated model and are sampled to a lesser extent in the higher protonation systems (Figure 3a,b). Stacking between wing II and the pyrimidine ring is clearly present in all simulations and is hypothesized to be relevant to the drug aggregate surface structure at the water interface. Analysis of ring-to-ring distance distributions at 0% protonation for core and surface molecules confirms that the internal structure of the aggregate displays crystal ordering while the surface is more disordered or liquid-like (Figure 3d).

**Surface Properties of Drug Aggregate Models.** Conformational and surface area analysis of compound **1** molecules at the aggregate/water interface suggests that torsional flexibility allows **1** to adopt surfactant-like orientations by exposing preferentially their “polar head” and burying their “hydrophobic tail”. We have categorized observed drug conformations in MD trajectories into four main groups: U,  $L_{\tau_3}$ ,  $L_{\tau_4}$ , and E (see Materials and Methods for details, Figure 1). The probability distributions for the  $\tau_3$  and  $\tau_4$  dihedral angles for core, surface, and dissociated molecules (Figure 4b) consistently indicate that the occurrence of partially and/or fully extended compound **1** conformations

( $L_{\tau_3}$ ,  $L_{\tau_4}$ , and E) correlates with a higher fraction of protonated drug molecules in the system. U conformations are predominant in the core of the aggregates in all simulations. Neutral compound **1** molecules at the surface in 10 and 30% protonated models show preference for  $L_{\tau_3}$  conformations (35.9 and 51.1% probability, respectively, see Table 1), while protonated compound **1** molecules at the surface of the aggregates prefer the U conformation with probability of 63.5 and 58.9%, respectively. Therefore, we conclude that protonated **1** molecules disfavor sampling of the  $L_{\tau_3}/L_{\tau_4}/E$  conformations while neutral molecules observe multiple forms. These observations highlight differences between the behavior of neutral and charged molecules on the aggregate surface.

In addition to looking at the drug aggregate packing motifs and conformational preferences, we also evaluated the makeup of the aggregate surface in terms of individual compound **1** atoms. Atom hydration analysis based on average normalized SASA of each heavy atom (see Materials and Methods, Figure 5a) show, as expected, that core molecules of **1** contribute little to the aggregate surface; however those core molecules that present atoms at the surface tend to display either cyano groups or pyrimidine rings. On the other hand, surface molecules, both neutral and protonated, prefer to expose wing II and pyrimidine rings. Hydration of neutral molecules at the surface is fairly uniform (Figure 5a) with preferences toward exposure of the pyrimidine ring for the 0% model and the cyano group in the



**Figure 4.** Compound **1** conformations and their occurrences in 0, 10, and 30% protonated model simulations. (a) Representative conformations of **1** observed in simulation: U conformation ( $\tau_3$  and  $\tau_4$  values between  $-90^\circ$  and  $90^\circ$ );  $L_{\tau_3}$  conformation ( $-90^\circ < \tau_4 < 90^\circ$  and  $90^\circ < \tau_3 < -90^\circ$ ).  $L_{\tau_4}$  conformation ( $-90^\circ < \tau_3 < 90^\circ$  and  $90^\circ < \tau_4 < -90^\circ$ ). E conformation ( $\tau_3 > 90^\circ$  and  $\tau_4 < -90^\circ$ ). (b) Probability distributions of compound **1** torsional angles  $\tau_3$  (A) and  $\tau_4$  (B) from the 0, 10, and 30% protonation models. Color assignment: light blue, all molecules in the simulation; dark blue, core molecules; magenta, surface molecules; green, dissociated molecules.

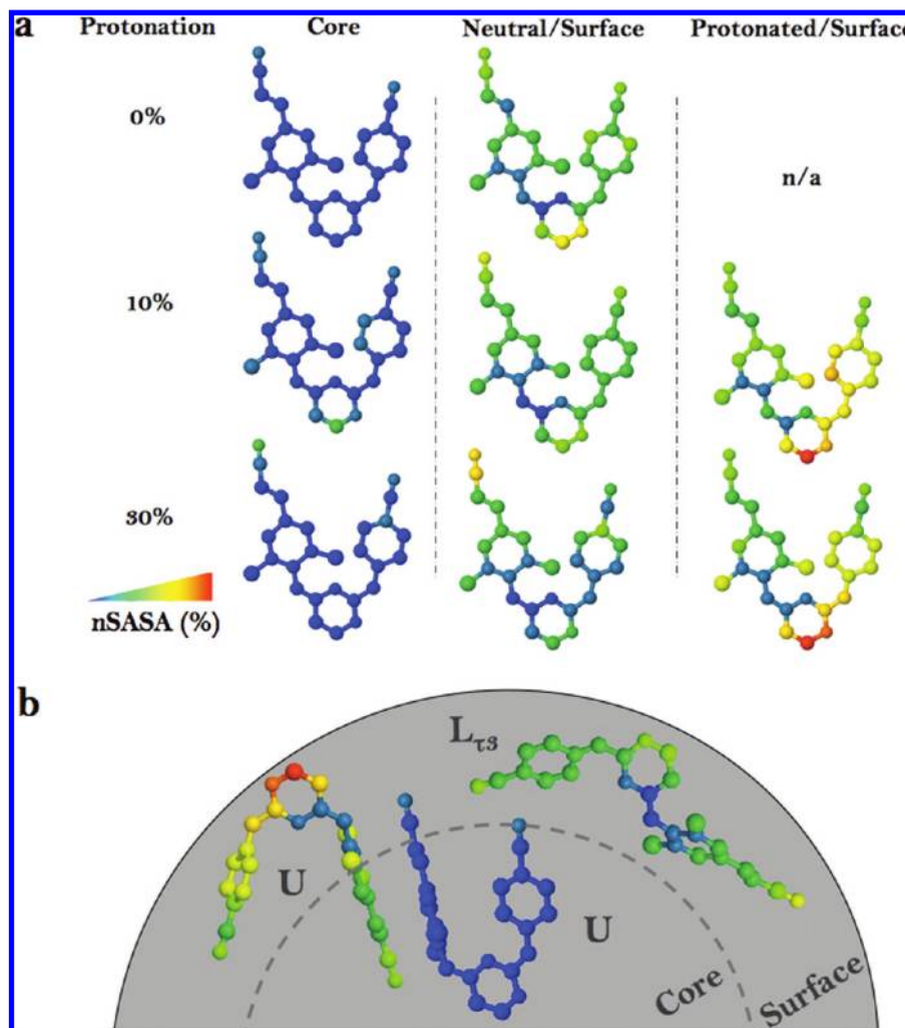
30% model. Charged surface molecules preferentially expose their pyrimidine rings at the surface of the aggregate with greatest hydration of the C(3) atom, shown in red (Figure 5a). On the basis of these results, we conclude that the contribution of core molecules to the surface is largely limited to cyano groups while surface molecules mostly display cyano groups, wing II, and pyrimidine rings. These findings agree with modes of binding determined in structural studies of RT/compound **1** complexes, where it has been observed that the cyanovinyl group of the molecule is oriented toward the solvent through a hydrophobic tunnel while the pyrimidine ring is partially exposed to the solvent region and hydrophilic residues on the other side of the binding pocket.<sup>15</sup>

Taken together, these observations suggest that molecules of **1** can exhibit different conformational preferences based on their environment. In Figure 5b, we summarize our conclusions concerning the orientations of the compound **1** molecules at the water/aggregate interface. Core molecules in the aggregate preferred the U conformations, while neutral, surface molecules sampled multiple conformations with preference toward  $L_{\tau_3}$  conformations that allows for surface

exposure of both the pyrimidine ring and cyano groups. Protonated **1** molecules assumed mostly the U conformations, preferentially exposing the charged pyrimidine ring while burying the mostly hydrophobic wings.

## Discussion and Conclusions

Pharmacokinetic and biophysical studies of **1** suggest that one possible explanation for the exceptional oral bioavailability properties of the drug lies with its ability to form  $\sim 100$ – $120$  nm diameter, self-formulating NNRTI aggregates and their subsequent uptake into systemic circulation.<sup>21,23</sup> We hypothesized that formation of the monodispersed aggregate populations is a product of the intrinsic properties of the compound **1** molecule. We proposed that aggregates of **1**, by size and morphology, resemble large micellar structures and/or microemulsions that are formed in the absence of conventional surfactants. Therefore, we hypothesized that the drug molecule is able to display surfactant-like properties under physiologically relevant conditions, facilitating micelle or microemulsion-like formation without the aid of added surfactants. The computational studies reported here support our hypothesis and suggest that under acidic conditions,



**Figure 5.** Compound **1** surface contribution analysis and summary. (a) Compound **1** atomic average solvent-accessible surface contributions computed from the 0, 10, and 30% protonation model MD trajectories. (b) Schematic diagram summarizing the observed preferred conformations and corresponding location (core and surface) in the aggregate. The diagram shows, starting from the left, a protonated molecule at the surface of the aggregate, a neutral molecule in the core of the aggregate, and a neutral molecule at the surface of the aggregate. Color assignment is based on normalized SASA values ranging from minimum (blue) to maximum (red) hydration.

protonation of **1** changes its behavior toward a more surface active molecule with clear preference toward exposure of hydrophilic portion of the molecule and burring of the hydrophobic groups. The computer simulations also show a clear change in the aggregate size and shape and a transition from solid-like to liquid-like phase with increasing number of charged species in the simulations. The models also suggest that under solution conditions of the simulation, the aggregates form a microemulsion-like phase with a partially disordered internal structure.<sup>10,27</sup> The structural flexibility and the headgroup size of the surfactant molecules are known to be fundamentally important for micelle and interfacial surface formation.<sup>33–36</sup> Smaller size, higher flexibility, and small electrostatic repulsions produce larger micelle structures and/or tighter surfactant molecule packing at the oil/water interface.<sup>10,27,33,35</sup> We believe that intrinsic flexibility of the compound **1** molecule, accompanied by its ability to assume amphiphilic properties under protonated conditions, similarly allows for more advantageous packing of drug molecules at the water/aggregate interface.

Aggregation at physiologically relevant conditions has been observed for many drugs and drug candidates, however, good

bioavailability properties are not as frequently observed for these molecules. In the series of studies reported by Seidler, McGovern, Feng, and other colleagues in the Schoichet group, formation of aggregate structures by drug and drug-like molecules has been correlated with promiscuous enzyme inhibition<sup>37–41</sup> and false positive results in HTS screening for drug leads.<sup>42,43</sup> In the recently published work by Coan et al., the drug aggregate composition is also compared to that of micelles.<sup>44</sup> Using the drug concentration measurements and monomer-to-aggregate volume ratio calculations, the authors concluded that drug aggregates were densely packed. These findings support our hypothesis of micelle-like drug aggregate formation and justify the close packing of the drug aggregates observed in our simulations. Our results also suggest that the mechanism of aggregate formation might be shared between promiscuous aggregators and molecules discussed in our work. While many drug molecules might form aggregate structures, we speculate that only those that are able to form these structures in a surfactant-independent fashion under physiologically relevant conditions might be able to follow absorption pathways associated with improved oral bioavailability. Conversely, we believe that drug or drug candidate

molecules unable to display amphiphilic properties under physiologically relevant conditions would tend to be less capable of forming stable, homogeneous populations of nanosize aggregates.

## Materials and Methods

**Dynamic Light Scattering (DLS).** Data collection and sample preparation were performed according to an earlier described method published in Frenkel et al. 2005.<sup>23</sup> Data collection was executed using a DynaPro MS800 at 25 °C. Each measurement consisted of 20 independent 10 s readings (repeated in triplicate). Data processing was done using DYNAMICS V6.7.60 software. Stock solutions of **1** were prepared by dissolving dry compound (>99% pure w/w, verified by reverse-phase chromatography) in 100% DMSO to a final drug concentration of 20 mM, filtered with a 0.22  $\mu\text{m}$  MillexGV sterile filter, and stored at -80 °C. Experimental samples were prepared by addition of a 20 mM stock solution of **1** to buffer at 1:199 volume ratio, followed by vortexing for 10 s. Buffer solutions were designed to mimic GI conditions of a fasting individual and spanned the pH range from 1.5 to 6.5. Buffers used in the study included the following: (1) 0.15 M HCl pH 1.5, (2) 20 mM malonate buffer pH 2.5 through 4.5, (3) 50 mM malonate buffer pH 5.5 and 6.5, (4) 20 mM malonate buffer pH 2.5 through 4.5 with 0.1% tyloxapol, and (5) 50 mM malonate buffer pH 5.5 and 6.5 with 0.1% tyloxapol.

Malonate buffers were prepared using malonic acid and sodium malonate solutions (J.T. Baker, Inc.). Tyloxapol (Sigma-Aldrich), a nonionic surfactant from the Triton group, was used to evaluate detergent-dependent aggregation behavior of **1**. Tyloxapol micelles with characteristic 3.5 nm hydrodynamic radii<sup>46</sup> were observed by DLS in control solutions and were clearly distinguished in experimental conditions from drug containing aggregate populations. Compound **1** was kindly provided by Janssen Pharmaceutica NV (Beerse, Belgium).

**Molecular Simulations.** The five models we have investigated differ in the fraction of protonated compound **1** molecules that was varied from 0 to 90% (see Table 1). The starting aggregate models were generated using small-molecule crystal structure coordinates of **1** in an anhydrous crystal form.<sup>45</sup> One hundred molecules of **1** were packed in the lattice according to the  $P2_1/c$  space group symmetry of the crystals with final microcrystalline dimensions of approximately 35 Å  $\times$  35 Å  $\times$  45 Å. These models were inserted using the Maestro Program (Schrodinger, LLC) into pre-equilibrated simulation cells of 60 Å  $\times$  60 Å  $\times$  60 Å containing pre-equilibrated water system (SPC water model). Water molecules that overlapped with the solutes were removed. In the resulting model, at least two layers of water molecules surrounding the generated drug crystals (see details in Table 1 and Figure 2). Depending on the model, a fraction of compound **1** molecules in the microcrystal were protonated at the N(2) atom of the molecule. The preferred protonation site at N(2) atom of the molecule was indicated by a recently obtained small-molecule crystal structure of protonated compound **1**.<sup>45</sup> Initial placement of protonated compound **1** molecules was performed in such a way to uniformly cover the surface of the microcrystal. If protonated drug molecules were present in the simulation, an equivalent number of chloride ions were included in the simulation to mirror observed solvent content in the small-molecule structure of the protonated compound **1** (Figure 2).

MD trajectories were obtained using the IMPACT molecular simulation program<sup>47</sup> with the 2005 implementation of the OPLS-2005 force field<sup>48-53</sup> (Schrodinger, LLC) under periodic boundary conditions and constant temperature and pressure at 298.15 K and 1 atm.<sup>47,54</sup> All covalent bonds were constrained using the SHAKE algorithm, with a relative tolerance of  $10^{-7}$ . The MD time-step was set to 2 fs. Full periodic boundary conditions were employed, and the molecule-based truncation was set at 10 Å distance for nonbonded interactions. Models

were equilibrated under constant isotropic pressure and minor changes to the volume were observed relative to the starting volume. Production runs were conducted at constant volume. The 60 and 90% protonated models were simulated for 7 ns and the 0, 10, and 30% protonated models for 9 ns. System coordinates were saved every 20 ps after a 2 ns initial system equilibration period. The resulting trajectories were analyzed using the Maestro program and associated utilities.

**Aggregate Cluster Analysis.** A molecule was considered part of the aggregate if it made significantly fewer water contacts than fully solvated molecules. To this end, we computed for each molecule  $m$  in each MD snapshot a solvation parameter  $S_m$  defined as the number of water molecules within 4 Å radius of any atom of the molecule. We defined a fully buried molecule in the core of the aggregate as having  $S_m = 0$ .  $S_m$  values greater than 20 correspond to fully dissociated drug molecules;  $9 < S_m < 20$  correspond to molecules on the surface of an aggregate, and  $0 < S_m < 10$  correspond to molecules that compose the interface between the surface layer and the core of an aggregate (Table 1). The aggregate size is expressed in terms of average number of molecules in an aggregate ( $N_{\text{agg}}$ ), and average number of surface, core, and charged molecules as reported in Table 1.

**Aggregate Surface Characterization.** MD trajectories were evaluated in terms of aggregate morphology, compound **1** torsional flexibility, aggregate packing, and drug molecule surface contribution as a function of drug location within an aggregate. Evaluation of aggregate morphology was done by visual inspection using the Maestro graphical interface in combination with solvent accessible surface area (SASA) estimation based on the periodic boundary-related contacts between models, if such were present, using a 1.4 Å radius probe.

To study behavior of **1** in the aggregate, we have categorized observed conformations into four main groups based on rotations of five dihedral angles. Torsion angles,  $\tau_1$ – $\tau_5$ , were identified using previously established nomenclature (Figure 1a). Drug conformations with  $\tau_3$  and  $\tau_4$  less than 90° are called “U” conformations (Figure 4a) and had been previously observed in crystallographic structures of the drug in protein complexes (wild-type and mutant)<sup>15</sup> as well as in the small-molecule crystal structures.<sup>45</sup> Drug conformations with  $\tau_3$  greater and  $\tau_4$  less than 90° result in rotation of wing I relative to pyrimidine ring and are called “L <sub>$\tau_3$</sub> ”, while those with  $\tau_3$  less and  $\tau_4$  greater than 90°, where wing II is rotated relative to pyrimidine ring, are called “L <sub>$\tau_4$</sub> ” (Figure 4a). The most torsionally flexed conformation of **1** is called “E,” for extended, and is characterized by  $\tau_3$  and  $\tau_4$  being greater than 90°, where both wings are rotated relative to pyrimidine (Figure 4a). The E conformation has been previously observed for a diaryltriazine analogue of a DAPY NNRTI molecule, R120393, in complex with HIV-1 RT and in a small-molecule crystal structure and has been called a seahorse conformation.<sup>45,55</sup>

The packing of the molecules in an aggregate was evaluated using ring-to-ring distance distribution analysis for 0, 10, and 30% models based on the distances between the centers of mass for selected ring pairs, sampling distances within a 10 Å radius. Both intra- and intermolecular ring-to-ring distances were calculated. On the basis of stacking interactions of the crystal packing observed in the anhydrous crystalline structure of **1**, we selected ring pairs for monitoring disruption of crystal packing as a function of protonation (Figures 3, 5). Distances of  $3.5 \pm 0.5$  Å correspond to intermolecular stacking interactions between wing I and wing II and the symmetry-related wing II rings in the compound **1** crystal packing.

The contribution of each atom to the aggregate SASA was computed with SURFV software using a 1.4 Å rolling probe.<sup>56,57</sup> Per atom surface contributions were normalized against the maximum SASA (mSASA) of an atom based on its most solvent exposed compound **1** conformation. Average SASA and number of surface charged molecules were used to estimate average aggregate surface charge density, reported in

Table 1. Figure 5 pertains to results of the hydration analysis, where the color gradient ranging from dark blue corresponds to small percent contributions to the surface and red (highest solvation) is used to accent the least and the most hydrated atoms of the molecules contributing to aggregate surface.

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