Adhesion of Photon-Driven Molecular Motors to Surfaces via 1,3-Dipolar Cycloadditions: Effect of Interfacial Interactions on Molecular Motion

Gregory T. Carroll,* Gábor London,* Tatiana Fernández Landaluce,† Petra Rudolf,‡ and Ben L. Feringa*,†,‡

†Centre for Systems Chemistry, Stratingh Institute for Chemistry, University of Groningen, Nijenborgh 4, 9747AG Groningen, The Netherlands, and ‡Zernike Institute for Advanced Materials, University of Groningen, Nijenborgh 4, 9747AG Groningen, The Netherlands

ABSTRACT We report the attachment of altitudinal light-driven molecular motors to surfaces using 1,3-dipolar cycloaddition reactions. Molecular motors were designed containing azide or alkyne groups for attachment to alkyne- or azide-modified surfaces. Surface attachment was characterized by UV—vis, IR, XPS, and ellipsometry measurements. Surface-bound motors were found to undergo photochemical and thermal isomerizations consistent with unidirectional rotation in solution. Confinement at a surface was found to reduce the rate of the thermal isomerization process. The rate of thermal isomerization was also dependent on the surface coverage of the motors. In solution, changes in the UV—vis signal that accompany thermal isomerization can be fit with a single monoexponential decay. In contrast, thermal isomerization of the surface-bound motors does not follow a single monoexponential decay and was found to fit a biexponential decay. Both one- and two-legged motors were attached to surfaces. The kinetics of thermal isomerization was not affected by the valency of attachment, indicating that the changes in kinetics from solution to surface systems are related to interactions between the surface-bound motors.

KEYWORDS: molecular motors · surfaces · photochemistry · click chemistry · adhesion

The use of organic chemistry to create molecular and supramolecular systems that undergo switching processes has inspired the design of numerous systems that can be cycled between states through the application of various stimuli.1–10 A key theme has been the construction of chemical systems that mimic dynamic functions of macroscopic machines with the ultimate goal of creating molecules that can perform useful work.11 The utility of rotary motions in macroscopic and biological machines provides a strong motivation for studying nanoscale constructs containing molecular components that can undergo rotary motions.12–14 Such systems are interesting not only in terms of understanding fundamental aspects of motion at the molecular level but also because biological systems have shown that rotary motions can be used to perform important functions. Two key examples are the ATP synthase15–18 and the bacterial flagellum,19 both of which use rotary motion for biological needs. These natural machines show that nanoscale rotary motors not only are interesting for academic purposes but also indicate their potential future role in powering artificial nanomachines.20 The potential for developing new technologies that utilize rotary motion has provided a strong impetus for understanding the physics of these biological motors as well as developing synthetic rotors12–14,21–32 whose chemical composition and physical properties can be tailored for a given device. In order to harness the collective motions of rotary motors to perform useful tasks, it is expected that surface-confined systems30,33–36 will be the most relevant, as these provide the opportunity to gain more order compared to solution-phase systems and minimize Brownian motion.37 In this report we study assemblies of altitudinal motors that can undergo controlled rotary motions (Figure 1).

We previously reported the synthesis and surface reactivity of the altitudinal light-driven molecular motor M1, shown in Figure 2.33 When attached to a surface, the motor retains its ability to undergo photochemical and thermal isomerization processes, reflecting the net unidirectional rotary behavior found in solution. Covalent attachment of the alkyne moieties of M1 to an interface likely forces the system to rotate in an altitudinal manner with the axis approximately parallel to the surface, affording opportunities in attempting to control the coordinates of adsorbates. Upon absorption of a photon, the central overcrowded alkene undergoes geometric
isomerization (Figure 2). The ground-state product of the photochemical transformation is less stable than the original molecule due to a change in helicity and orientation of the stereocenter. The methyl substituent at the stereogenic center adopts the less stable pseudoequatorial orientation instead of the more stable pseudaxial orientation. In order to relieve strain, the motor undergoes an irreversible thermal helix inversion. During the thermal isomerization the methyl substituent at the stereogenic center changes from a pseudoequatorial to a pseudoaxial orientation. A second photochemical and thermal isomerization completes the rotary cycle. Although much work has been done on these systems in regard to controlling the speed of rotation in solution, macroscopic surface-attached systems are largely considered to be more practical in converting rotary motion into mechanical work. The current paradigm regarding the thermal isomerization takes into account only the internal structure of the motor; yet key applications of these molecules are expected to involve interactions beyond steric hindrance within a single molecule. In this report we investigate the importance of confinement on the speed of rotation. We show that intermolecular interactions significantly affect the rate of thermal isomerization, a key step in the rotary cycle that ensures unidirectional rotation. The results are particularly significant with respect to key challenges in the area of prototypical nanomachinery: many molecular motors working in concert and the use of rotary molecular motors to induce the mechanical motion of an adsorbate. Additionally, the results indicate that a rotor of the kind described herein can undergo rotary motion in a confined environment in which intermolecular interactions impede thermal isomerization. Although our motivations are related to the construction of nanoscale machinery that is powered by controlled molecular motions, the system described herein has general implications for surface interactions and their effects on molecular mechanics.

RESULTS AND DISCUSSION

As described previously, silanated surfaces (SAM 1) containing azides were prepared on quartz and silicon substrates (Figure 3). Immersion of substrates containing Si—OH surfaces in dilute solutions of silyl ethers is known to form stable coatings typically described as self-assembled monolayers (SAMs). We found that immersion of piranha-cleaned quartz or Si/SiO2 substrates in 1 mM solutions of compound 1 consistently gave organic coatings of 1.8 nm or less as measured by ellipsometry. The theoretical length of compound 1 is approximately 1.9 nm, consistent with a monolayer of the azide. A given substrate showed a thickness spread of no more than 0.1 nm when multiple areas of the substrate were measured, indicating a uniform coating. The coatings were stable to sonication in toluene and methanol. The most efficient procedure for achieving a high coverage of azide involved self-assembly from a 20:1 cyclohexane/THF solution containing a small quantity of H2O and HCl (see Supporting Information). Self-assembly in toluene reduced the coverage by approximately 50%. Self-assembly in cyclohexane without water or acid required two days to reach a film thickness of approximately 1.7 ± 0.1 nm. In addition to the
XPS and contact angle measurements previously performed on SAM 1, ATR-IR spectroscopy, performed by placing the surface in intimate contact with a germanium crystal, was used to show the presence of the azide functionality on the substrate (Figure 4). The distinct band at 2095 cm\(^{-1}\) clearly shows the presence of an azide on the substrate. Collectively, the characterization shows a highly stable and uniform coating of azide-functionalized molecules with a thickness that is consistent with a monolayer. As will be discussed below, the surface-bound azides provide a highly reactive nanolayer for attaching alkyne-modified molecules.

M1 was previously shown to adhere tenaciously to silanated surfaces containing an azide, attributed to covalent bond formation induced by a 1,3-dipolar cycloaddition (Figure 3). A variety of characterization techniques support the hypothesis that the motor adheres to the surface via covalent bond formation. Most simply, the UV–vis spectrum of a quartz substrate shows the presence of the motor after immersion in a 1 mM solution of M1 in the presence of copper catalyst. Note that the surface was rinsed by sonication for up to three minutes in DMF, H\(_2\)O, and MeOH (1 × each solvent). Immersion of an azide substrate in the solution without catalyst or immersion of a quartz slide without azide does not result in the adhesion of M1 at room temperature or with heating at temperatures of 65 °C, even after 5 days of immersion. This indicates that the signal is not simply the result of insufficiently rinsing a surface befouled by adsorption from solution. Rather, the silane coating is chemically modified by the controlled reaction of a carefully designed molecule. The amount of catalyst was varied from 1 to 5 mol % relative to the alkyne.

Figure 4. ATR FT-IR spectra of SAM 1 and MS1.
resulting UV signal was observed. Likewise, performing the surface reaction at 65 °C showed no difference in the intensity of the signal. Changing the immersion time provided a simple method to control the coverage. After removing the substrate at various times followed by sonication (2 min 3× in toluene and MeOH), the UV signal increased with reaction time, as expected for a covalent attachment to the interface (S1). The intensity of the UV signal and reaction time to reach saturation were dependent on the quality of the azide layer. A higher coverage of azide produced a higher UV signal and a longer reaction time to reach saturation. The higher density of motor was also evident in a higher ellipsometric thickness. The highest ellipsometric thickness of the motor layer obtained for the highest yield of azide is approximately 1.5 nm, which is close to the calculated length of 1.55 nm if the "legs" of M1 are oriented normal to the substrate. The UV–vis and ellipsometry measurements previously reported33 clearly indicate the presence of the motor on the azide-coated surfaces, which appears to be a single layer. Covalent attachment of the motor to the surface seems the likely adhesion mechanism given the control experiments described above. XPS measurements showed the disappearance of the azide signal and the presence of the triazole group, strongly supporting attachment through a 1,3-dipolar cycloaddition. To further support our claim, ATR-IR spectroscopy was performed on the surface (Figure 4). As noted above, the azide signal is clearly present on the silicon substrate. After reaction with the motor, the signal attenuates to the level of noise, providing further evidence for covalent modification of the azide layer. A reduction in the H2O contact angle from 82 ± 1° to 65 ± 1°, probably due to the ethylene glycol legs and triazole moiety of the surface linkage, was also previously used to characterize the coupling reaction and implies that the layer of motors is not so well packed as to screen the molecular components beneath the chromophore from a water droplet. Note that the H2O contact angle of a poly(styrene) film is approximately 90°.42 Although not a perfect comparison, it provides a comparative value for a smooth film composed of aromatic units and aliphatic CH groups, both of which comprise the motor component of M1. The combination of characterization techniques and resistance to repeated cycles of sonication and washing strongly indicates that the motors are covalently bound to the interface.

An alternative approach one may opt for is to directly assemble a silanated version of the motor, M2 (Figure S2). Although such an approach appears more efficient on paper, an advantage of the interfacial 1,3-dipolar cycloaddition of the motor rather than hydrolysis of a silanated motor is that multilayer formation during assembly or polymerization during storage or handling is no longer a possible undesirable outcome. Additionally, performing the reaction on the surface allows one to bypass a step of the synthesis and the accompanying purification methods. In any case, we felt that it would be instructive to work with a silanated version of the motor. Our first attempt at forming the triazole in solution and assembling the silane M2 resulted in a multilayer of 77 ± 8 nm and an H2O contact angle of 79 ± 3° on silicon. Polymerization of the silane was evident in the formation of a yellow particulate within ten minutes of mixing a THF solution containing silane and a small amount of H2O and acid in cyclohexane. Scanning electron microscopy (SEM) images show what appear to be fused particle-like aggregates (Figure S2). A stable coating was formed on silicon and quartz that was resistant to sonication. Irradiation of the film with UV light produced a yellow film that faded with time as the motor underwent thermal isomerization to the stable form. The changes in the UV–vis spectrum correspond to the changes found in solution and for the monolayer (Figure S2). Thinner films containing a thickness that corresponds to monolayer coverage could be obtained by reducing the concentration. For example, a film with an ellipsometric thickness of 1.9 ± 0.1 nm could be obtained by assembling from a 0.01 mM solution (S3). Interestingly, self-assembly from THF showed very little coverage (0.3 nm ellipsometric thickness) and may be a result of competition with THF coordinating to the surface, as has been implicated when self-assembly is attempted on mica43 (S3), although this comparison is imperfect, as mica is chemically different and may have a different mechanism of adhesion prevention. Although the system can be somewhat intratable, it provided an additional variant to study the kinetics of isomerization, as will be discussed below.

In order to more directly probe the feasibility of two-legged attachment as well as expand and test the versatility of the interfacial reaction, we synthesized motor M3, which contains two azides for attachment to an alkyne-modified surface, SAM 2 (Figure 5). We silanated quartz and silicon wafers with 6-(propargyloxy)-N-(triethoxysilylpropyl)urethane. After reaction of the film with M3 under the same conditions for attaching M1 to an azide surface, the UV–vis spectrum showed the presence of the motor on the surface only when catalyst was added to the reaction mixture (Figure S4). In the absence of catalyst or an alkyne film no motor was detected on the quartz surface. The distinctive IR and XPS signals corresponding to an azide film no motor was detected on the quartz surface. The distinctive IR and XPS signals corresponding to an azide functionality were used to probe the propensity for two-legged attachment.33,44 Both the IR and XPS spectra show no evidence for unreacted azide in strong support that both legs of the motor react under the conditions employed (Figure 6).

As previously shown, M1 can undergo photochemical isomerization followed by irreversible thermal isomerization while confined at an interface.33 The thermal isomerization is the rate-limiting step of the rotary cycle. Understanding how the structure of the motor af-
ffects the half-life of the thermal step has been key in increasing the speed of the motor. A general paradigm regarding the effect of substituents and ring size has been developed in order to design faster motors.\textsuperscript{38} Although much progress has been made in synthesizing motors of various speeds, confinement and intermolecular effects on the speed of rotation have previously received little attention. In solution the thermal isomerization of \textit{M1} was found to proceed through a unimolecular process and a half-life of 90 s in methanol. In order to compare the surface-bound system with the solution analogue, we followed the time dependence of the changes in the UV spectrum during thermal isomerization at the interface using a surface that had a coverage of approximately 50% (Figure 7). In contrast to the solution experiments, fitting the thermal decay
of the signal at 439 nm suggested at least two processes occurring during the thermal isomerization that have half-lives that differ by an order of magnitude at 25 °C. The first half-life had a value of approximately 110 s, not so different from the half-life of 90 s measured in MeOH solution. More striking is the extraction of a second half-life of 1200 s, making the rotary cycle of the surface-bound ensemble considerably slower than in solution. We postulate the existence of at least two different processes, as evidenced by the biexponential decay of the UV signal corresponding to the unstable form of the motor. The biexponential fit may reflect environmental or configurational heterogeneities (i.e., surface roughness, defect sites, etc.) or multiple processes occurring during thermal isomerization when M1 is clustered and bound to a substrate. As expected, higher temperatures increase the rate of thermal isomerization on the surface. For example, measurements performed at 40 °C gave a thermal decay for which half-lives of 30 and 270 s were obtained.

Interestingly, structurally different surface-confined systems have similarly shown a non-monoexponential decay corresponding to the kinetics of molecular conformational changes. For example, methylthiolate chemisorbed on a Ag(111) surface showed a biexponential decay for the lifetime of the first excited level of the symmetric C—H stretching mode. Similarly, thin polymer films containing azobenzenes attached to side-chains have shown thermal rates that deviate from simple monoexponential behavior. Regardless of the fitting procedure, it is clear that the thermal conformational change requires more time on the surface compared to solution. On the basis of our model of a layer of oriented motors covalently attached to a solid substrate, supramolecular interactions within a cluster of surface-bound motors seems to be the most plausible reason for the observed kinetics. The hypothesis was tested by constructing surfaces with a different coverage of motors and comparing the half-life for thermal isomerization. A potential advantage of the surface modification strategy presented herein is that better control over the density of the bound species can be obtained compared to direct self-assembly of a silane, which can assemble into dense clusters prior to surface attachment. Although the exact mechanism of interfacial 1,3-dipolar cycloadditions for small molecules is not known, we assume that for M1 self-assembly into clusters at the surface does not precede covalent bond formation. For our arguments below we will assume random growth of the film without preaggregation or self-assembly; however we cannot explicitly rule out the latter. In order to understand the effect of surface coverage on the kinetics of the thermal isomerization, we examined four different coverages. We designate our average on the kinetics of the thermal isomerization, we latter. In order to understand the effect of surface coverage on the kinetics of the thermal isomerization, we examined four different coverages. We designate our assembly; however we cannot explicitly rule out the dom growth of the film without preaggregation or self-assembly. For our arguments below we will assume ran-

Figure 7. Thermal decay of UV signal of M51 (50% surface coverage) at 439 nm at 25 and 40 °C. Both data sets were fit with a biexponential decay, and hence two half-lives for the rate of thermal isomerization were calculated for each temperature. The half-lives at 25 °C are 110 and 1200 s. The half-lives at 40 °C are 30 and 270 s.

approximately 50% and 25% the saturation coverage (saturation factor of 0.5 and 0.25, respectively). Additionally, we made use of our multilayer formed by assembly of the silane M2 in order to probe a more extreme situation in which a given motor is more susceptible to axial interactions compared to a monolayer in which axial interactions could occur only when deviations from our model in Figure 1 occur (i.e., surface roughness, defects that considerably tilt the axle, adventitious impurities). The half-lives for the thermal isomerization are presented in Figure 8 (see Figure S5 for thermal decay curves). Both the fast and slower processes show a decrease in half-life with a decrease in the surface coverage. At full coverage, the half-lives are 280 and 4100 s, approximately twice that found for surfaces with 50% coverage. The most drastic change occurs on going from the monomolecular layer to the multilayer prepared by polymerization of the silane. The overlayer experienced by the motors in the multilayer film has a significant effect on the motors, which not only provides support for the idea that supramolecular interactions affect the speed of the rotary cycle but also indicates that the systems fabricated by the controlled dipolar cycloaddition do contain a reasonable degree of order as expected from chemoselective covalent bond formation of a small molecule with a surface.

One may consider that the two processes indicated by the biexponential fit reflect surface populations that are attached through one and two legs. If this was the case, then attaching a one-legged motor would show a monoexponential decay. In pursuit of understanding the valency of surface attachment on the kinetic behavior of the rotary cycle, a one-legged derivative of M1, M4, was synthesized (Figure 9). The UV—vis absorption spectrum is similar to that obtained for M1 (Figure S6). The maximum of the longest wavelength absorption is blue-shifted by 3 nm in MeOH and 4 nm on the surface.
When 1 mol % of copper catalyst was used, M₁ was found to react with the surface much faster than M₄, as determined by the UV–vis spectrum. The UV–vis spectra reached saturation almost after 12 h for M₁, whereas the surface reaction for M₄ was only approximately halfway complete, requiring an additional day. As found for M₁, the azide peak in the IR spectrum disappeared (S7). Similarly, the azide band at 405 eV in the XPS spectrum is significantly attenuated (Figures S7, S8, S9, S10), signifying chemical transformation through the formation of a triazole moiety. The decay of the signal at 439 nm was monitored following irradiation with UV light under the same conditions employed for M₁ (Figure S5). As found for M₁, the resulting decay showed two processes, the first with a half-life of 240 s and the second with a half-life of 4500 s, similar to the values obtained for the two-legged system (Figure 8). The presence of two rates and the slower thermal isomerization indicate that the kinetic behavior of the rotary cycle is more strongly affected by factors other than one or two-legged attachment. The most plausible explanation is intermolecular interactions between the surface-bound motors. Regardless of the origin of the behavior, it seems clear that confinement of a group of motors at an interface or within a film considerably increases the half-life of the rotary cycle.

We have shown that the Cu-catalyzed 1,3-dipolar cycloaddition reaction provides a versatile approach to attach molecular motors to surfaces. When azide motors are attached to alkyne surfaces, the absence of both XPS and IR signals corresponding to the azide strongly suggests predominantly two-legged attachment. Attachment of motors to surfaces increases the half-life of the thermal isomerization. The speed of the motor increases when the coverage is lower and severely decreases when multilayer polymer films are formed. Additionally, motor films attached to quartz surfaces show at least two half-lives for thermal isomerization, pointing to significant intermolecular interactions between motors, affecting rotary speed. The incorporation of functional groups onto one side of the rotor will allow for more advanced interfaces to be constructed in which surface properties can be reversibly transformed and patterned with photons.41,47

Figure 8. The bar graphs present the half-life values for thermal isomerization as a function of surface coverage for both the one- and two-legged systems.

Figure 9. Surface attachment of one- and two-legged motors.
Acknowledgment. Financial support from NanoNed (G.L., G.T.C., T.F.L.), The Netherlands Organization for Scientific Research (NWO) (G.T.C., B.L.F.), Zernike Institute of Advanced Materials, the Materials Science Centre (MSC’), and the Foundation for Fundamental Research on Matter (FOM) (T.F.L., P.R.) is acknowledged.


42. Wu, S. Polymer Interface and Adhesion; Marcel Dekker: New York, 1982.


