In principle, excitation of surface plasmons by molecular tunnel junctions can be controlled at the molecular level. Stable electrical excitation sources of surface plasmons are therefore desirable. Herein, molecular junctions are reported where tunneling charge carriers excite surface plasmons in the gold bottom electrodes via inelastic tunneling and it is shown that the intermittent light emission (blinking) originates from conformational dynamics of the molecules. The blinking rates, in turn, are controlled by changing the rigidity of the molecular backbone. Power spectral density analysis shows that molecular junctions with flexible aliphatic molecules blink, while junctions with rigid aromatic molecules do not.

1. Introduction

Surface plasmon polaritons (SPPs) can carry optical information at petahertz frequencies along sub-diffractional metallic waveguides, which enables new designs for integrated optoelectronic devices.[1–3] Although the optical properties of the plasmonic structures, such as metallic particles, meta-surfaces, and waveguides, have been extensively studied,[4–7] there is still a lack of robust on-chip plasmon sources that convert electrical signals to plasmonic signals.[8–12] To overcome this challenge, traditional light sources, such as light-emitting diodes (LEDs), can be miniaturized and integrated with plasmonic waveguides.[8–12] Another approach is to use tunnel junctions as on-chip plasmon sources, that directly excite plasmons by electrons that tunnel inelastically.[13–17] In the former approach, junctions are excited in two steps involving a relatively slow electron-hole recombination process (>ps),[18] while in the latter approach plasmons are excited in a single step at much faster quantum tunneling timescales (=fs).[15,17,19]

Recently, we have demonstrated on-chip electrically driven plasmon sources using self-assembled monolayer (SAM)-based tunnel junctions (STJs) and showed the control over the plasmonic properties (such as intensity, polarization, and frequency) by the chemistry of the molecules in the STJs.[20] Under applied bias, the STJs excite surface plasmons from a discrete number of diffraction-limited spots, which blink between “on” (bright) and “off” (dark) states with on- (\(t_{\text{on}}\)) and off- (\(t_{\text{off}}\)) probability densities ranging from microseconds to minutes. We characterized the on-time \(P(t_{\text{on}})\) and off-time \(P(t_{\text{off}})\) probability densities of the intermittent light emission (blinking) in the STJs and found that the \(P(t_{\text{on}})\) and \(P(t_{\text{off}})\) probability densities follow a power-law dependence similar to other point-like sources, such as single chromophores or quantum dots.[21] Here, we reveal the underlying mechanism that causes blinking in STJs and how it relates to the conformational changes of molecules in the junction. We use this insight to control the intermittency of the light emission by changing the molecular backbone rigidity (defined as the amount of energy required to rotate the molecular backbone around the C-C bond, see Supporting Information and also below). These findings are important for the field of molecular electronics as it is explicitly demonstrated here that the dynamic aspects of the molecule in-between the electrode interfaces must be taken into account in device engineering.[22–28]

In molecular electronics, a challenge is to rationalize the large spread of eight orders of magnitude in the measured values of \(J\) (the current density) for the same molecule incorporated in different types of junctions.[28–37] Although it has been proposed that this discrepancy can be (largely) explained by assuming that the effective electrical contact area between the SAM and the top electrode is orders of magnitude smaller than the geometrical
contact area, reliable methods to measure the effective electrical contact area are not available. It is well-known that the effective electrical contact area between two solid surfaces can be several orders of magnitude smaller than the geometrical contact area due to the surface roughness of the electrodes, in molecular electronics, however, it is common practice to assume that all molecules participate in charge transport and that the top-electrode always makes good contact to the molecules regardless of their chemical nature. In addition, the dynamics of molecule–electrode contacts are usually ignored and assumed to be static, while molecules have of course conformational degrees of freedom depending on their chemical structure.

We believe that the intermittent light emission of molecular electronic plasmon (MEP) sources can be explained by the fluctuating barrier model associated with the conformational changes of the molecules in response to inelastic tunneling where charge carriers interact with the vibrational modes of the molecule resulting in conformational changes of the molecules. As a result of these molecular conformational changes, the effective tunneling barrier width and height fluctuate resulting in current fluctuations. The dynamics of the molecular conformational changes has been well-studied in molecular tunnel junctions and followed in real-time by recording the change of tunneling current as a function of time.

Zandvliet et al. demonstrated convincingly that inelastic excitation-induced conformational changes can be used to trap single molecules between a scanning tunneling microscope (STM) tip and a surface.

Under continuous excitation, point sources, such as quantum dots and single molecules, emit light irregularly, and often switch between "on" and "off" states with the times and that characterize the length of each state ranging from microseconds to minutes. The and follow an inverse power law, where and range from 1 to 2. Another important feature of blinking is the power-law dependency of the power density where is the frequency domain and is the Fourier transform of the autocorrelation of the emission intensity time trace . Similar to , the distributions of follow a power law in the form of where is between 1 and 2. The behavior is often modeled with Equation (1)

\[
S(f) = \frac{A}{f} + \frac{B}{(f/f_0)^\alpha} + C
\]

where and are the amplitudes of the 1/f (also called flicker noise) and the 1/f^2 components, is the characteristic frequency of the 1/f^2 component and can be characterized by the inflection point in S(f), and the constant does not depend on frequency and is an offset representing the white noise floor.

Equation (1) is widely used to evaluate the fluctuations in time traces, and has been used to understand the intermittent light emission of point sources.

In this work, we experimentally demonstrate the control of the intermittent light emission of the MEP sources by the rigidity of the molecular backbone. With stiff aromatic molecular backbones, the blinking is almost eliminated, while the opposite is true for flexible aliphatic molecular backbones over the range of frequencies (10^1–10^9 Hz) studied here. These observations suggest that conformational changes of the molecules play a major role in the blinking of MEP sources. This work also highlights that the electrical contact between the electrode and the barrier molecules is highly heterogeneous and varies from junction to junction depending on the chemical nature of the molecules. In addition, the electrical contact points along which tunneling occurs change within the junctions resulting in time-dependent changes in the junction properties.

2. Results and Discussion

Figure 1 shows the two types of STJs with two different types of molecular backbones. We used STJs with SAMs of dodecane-1-thiolate (in short SC12, Figure 1a) and (4-(phenylethynyl)phenyl)methanethiolate (in short SC-PEP, Figure 1b). Because of the sp^2 character, the SC-PEP molecules have a more rigid molecular backbone than the sp^3 aliphatic SC12 molecules.

Following the same methods (see the Supporting Information) as described by Du et al., we prepared the two types of STJs using semitransparent 50 nm gold bottom electrodes obtained by template-stripping onto which we formed the SAMs. The top electrodes were formed using EGaIn (a eutectic gallium-indium alloy with 0.7 nm native Ga_2O_3), which is well-established and generates high-quality molecular junctions in high yields of working devices.

EGaIn is a non-Newtonian liquid metal that flows when shear pressure is applied. When brought in contact with the SAM, EGaIn flows and deforms for an instant, before solidifying again. This feature of EGaIn ensures that a soft top-contact is generated and that the SAMs retain their supramolecular structure, essential to obtain a good tunneling barrier. In our experiments, the EGaIn was confined in a through-hole in a polydimethylsiloxane microfluidic device and the top-electrode had a geometrical contact area of 1000 µm^2.

Figure 1 also shows schematically the difference in the conformational degree of freedom between SC12, which, e.g., can have gauche conformations which are not possible for SC-PEP. Consequently, an SC12 molecule can be in contact with both electrodes when in an all-trans conformation but the contact with the top-electrode will be lost when a Gauche defect is present. In the former case, the tunneling barrier width (d_t) and height and the corresponding tunneling decay coefficient (β_t) are defined by the molecule (solid line in Figure 1c). In the latter case, a gap between the molecule and the top-electrode is present and, consequently, besides the molecular component (defined by d_i and β_i) a second component is presently defined by the air gap d_a and associated tunneling decay coefficient β_a (dashed line in Figure 1c). Since through-molecular bond tunneling is more efficient than tunneling through air, the local tunneling current decreases significantly along the molecule when an air gap forms. Hence, the local current increases elsewhere inside the junction until the molecule changes its conformation again since the total current across the junction remains the same with variations within noise levels (see previous works for J(t) measurements). Therefore, when the...
current across the molecule is low, the plasmon excitation rate (and thus the photon emission rate) is low, and vice versa. In other words, this conformational change (and other types of conformational changes) results in a fluctuating tunneling barrier and consequently in intermittent light emission. Although SC-PEP molecules are “stiff,” conformational changes can still lead to current fluctuations due to, for instance, rotation of the phenyl rings (Figure 1d). It is well known that when the two phenyl groups are out of plane (large dihedral angles), the conductance along the molecule can be up to ten times smaller than when the phenyl groups are in plane (small dihedral angles). \[^{61,62}\] In the SAM, however, these rotations are hampered. \[^{63,64}\]

The \(J(V)\) characteristics of the two types of STJs were recorded using the Keithley 6430 source meter and a homemade Labview program, with electrical bias applied between \(-1.8\) and \(+1.8\) V in steps of 90 mV. Figure 2 shows the log-average \(J(V)\) curves (averaged over 150 traces) of the two types of STJs. We note that the two SAMs have a similar molecular length (1.8 nm for SC\(_{12}\) and 1.6 nm for SC-PEP), thus junctions with SC-PEP SAMs with the aromatic molecular backbone, in theory, should have higher \(J\) values than junctions with SC\(_{12}\) SAMs because aromatic molecules have smaller tunneling decay coefficients (0.2–0.4 Å\(^{-1}\)) than aliphatic molecules (0.8 Å\(^{-1}\)) \[^{32–36}\] see Equation (2) below. This discrepancy can be explained by the difference in the effective electrical contact area which is directly visualized in the light emission images from the two types of STJs shown in Figure 3 (and see below for more details).

To record the light emission from the STJs in real-time, we used a wide-field inverted optical microscope equipped with an electron-multiplying charge coupled device and a 100 \(\times\) oil immersion objective with the numerical aperture \(NA = 1.49\) (see the Supporting Information). Figure 3 shows real plane emission images of the two STJs recorded over a period of time of 120 s (1 s per image) at \(-1.8\), \(-2.0\), and \(-2.2\) V bias (see also Figures S1–S3, Supporting Information for additional light emission images and spectra). By comparing Figure 3a,b, we make the following observations: i) the emission spots in STJs with SC-PEP SAMs blink less frequently than STJs with SC\(_{12}\) SAM (see also Figure 4a,b and Supporting Movies, Supporting Information), ii) the blinking rate increases with increasing.
the molecules is much larger (4.5 eV for SC-PEP[65] and 8–9 eV molecular orbital–lowest unoccupied molecular orbital gap of electroluminescence) of the molecules as the highest occupied molecular orbital–lowest unoccupied molecular orbital gap of the area with no emission cannot be caused by “quenched behavior” of the molecules that keep their conformations stable over prolonged periods of time up to hours.

To quantify the blinking dynamics, we performed $S(f)$ analysis of the intermittent light emission from individual emission spots for both types of STJs (see Figure S5, Supporting Information). Figure 4 shows representative time traces of the light emission intensity recorded over 1000 s at an applied bias of $-2.0$ V along with the $S(f)$ analysis. The $I(t)$ of the STJ with SC$_{12}$ SAMs (Figure 4a) shows clear jumps (>10k counts) between the on and off states. However, the emission of the STJ with SC-PEP SAMs (Figure 4b) remains in the “on” state and displays small variations in intensity (<5k counts). Within the experimental frequency range ($10^{-3}$–1 Hz), the $S(f)$ of the SC$_{12}$ junction (Figure 4c) decreases with increasing $f$ and has a maximum value of around 0.1 Hz. The solid red line is a fit to Equation (1) with three components: the 1/$f$ component at the low frequencies, the 1/$f^2$ component with $f_0 = 0.1$ Hz, and the white noise component (at the level of $2 \times 10^{-4}$ a.u.). In contrast, the $S(f)$ of the SC-PEP junction (Figure 4d) decreases as a function of $f$ and only has the 1/$f$ and the white noise components (at the level of $1 \times 10^{-4}$ a.u.). Here, the 1/$f$ component results from intensity fluctuations that occur randomly in the on state and do not have a measurable repetition period. While in the case of SC$_{12}$ junctions, the 1/$f$ component is mainly due to the random intensity fluctuations of the off states. The extra 1/$f^2$ component with the characteristic frequency $f_0 = 0.1$ Hz corresponds to the pronounced intensity fluctuations (the blinking) between the on and off states with a repetition period of around 10 s. It is important to note that although the time scale for the molecular conformational change can be very fast, the conformational changes in SAM are very slow due to steric hindrance between neighboring molecules and favorable packing energies. Previous STM experiments[50,68,69] show that the alkanethiol and oligo(phenyleneethynylene) molecules in SAMs keep their conformations stable over prolonged periods of time up to hours.

It has been reported before that the rate of conformational changes induced by inelastic tunneling increases with increasing applied bias.[42–50] Therefore, we studied the bias dependency of the blinking behavior by determining $S(f)$ of the SC$_{12}$ junctions at $-1.8$, $-2.0$, and $-2.2$ V. Figure 5 shows representative time traces of the light emission intensity along with the $S(f)$ analysis. For each applied bias, we analyzed 40–50 individual blinking spots to determine $f_0$ and Figure 5c shows the histograms of $f_0$ with a Gaussian fit (because the blinking of the emission spots is not correlated, but random) to these histograms to determine the Gaussian mean value of $f_0$. The value of $f_0$ increases from 0.09 Hz at $-1.8$ V to 0.15 Hz at $-2.2$ V.

Figure 2. $J(V)$ curves of STJs with a) SC$_{12}$ and b) SC-PEP SAMs in the bias range of ±1.8 V. The $J(V)$ values are averaged over eight junctions, and the error bars are the corresponding log standard deviations determined over 150 $J(V)$ traces.

The current is much larger for the SC$_{12}$ junctions compared to SC-PEP junctions due to the poly-crystalline and non-Newtonian properties of the GaO$_x$/EGaIn contact resistance.[31,38] The contact resistance is mainly due to the non-Newtonian properties of the GaO$_x$/EGaIn interface, which reduces the effective contact area further explaining the low effective contact areas we report here.[67]
showing that the blinking rate increases with bias. These observations agree with inelastic tunneling: with increasing bias both the tunneling electron energy and the tunneling electron rate increase, thus the possibility for the conformational change increases, which explains the near doubling of \( f_0 \). For SC-PEP junctions, we also observed an increase in the blinking rate with increasing bias, however, the \( 1/f \) component dominates in the measured bias range (see Figure S4, Supporting Information).

To investigate how the tunneling current influences the blinking of the light emission in STJs, we also performed the \( S(f) \) analysis of the time traces of the tunneling current. Figure 6 shows the tunnel current versus time traces of the STJs with SC\(_{12}\) SAMs and SC-PEP SAMs with the current sampling time 0.13 s (3.8 times faster than the optical case, 0.5 s). For both types of STJs, the tunneling current fluctuations are smaller than 20% and only show the \( 1/f \) component. This cannot account for the \( 1/f^2 \) blinking behavior of the SC\(_{12}\) junction. We note that the recorded tunneling current represents the current flowing across all spots in the entire STJ. As shown in Figure 3 and the Supporting Videos (Supporting Information), the blinking of the emission spots is not correlated. Consequently, the local current fluctuations are also not correlated and therefore current fluctuations at local spots cancel each other within the noise level of the measured total currents (see Figure S6, Supporting Information). Thus, the advantage of the optical characterization becomes obvious as it allows for independent analysis of each individual light-emitting spot.
providing local information on how the current flows across an individual spot at any given time. We also note that, in tunnel current time trace measurements, the term $C$ represents the thermal noise of the tunnel junctions. For both types of junctions, the current is at the same level (within a factor 2–4) and the term $C$ of $5 \times 10^{-5}$ is also very similar (Figure 6c,d).

We estimated the size of the air gap required to induce a significant change of the local current for the SC$_{12}$ molecule (Figure 1c) as follows. The value of $J$ can be described by the general tunneling equation (Equation (2))

$$J = J_0 e^{-\beta_2 d_2}$$

(2)

The tunnel barrier can be described as a double barrier junction by Equation (3) with a gap of air

$$J' = J_0 e^{-\beta_2 d_2} e^{-\beta_1 d_1} = J_0 e^{-(\beta_1 + \beta_2) d_2} e^{-\beta_1 d_1}$$

(3)

where $\beta_2$ is the tunneling decay coefficient of air (since our experiments are conducted in ambient conditions). Since the value of $\beta_2$ of air is 2.9 Å$^{-1}$ and $\beta_1$ is 0.8 Å$^{-1}$ for aliphatic SAMs,[30,34–36,38] a value of $d_2$ of 1.1 Å (which is roughly the length of one CH$_2$ unit) results in a factor of 10 decrease in the value of $J$. Such a small gap can be easily obtained by bending of the molecule and (partial) rotation of molecular segments around C-C bonds. For aromatic SAMs, $\beta_1$ is 0.2–0.4 Å$^{-1}$,[12–36] and a value of $d_1$ of 0.9 Å is sufficient to reduce the local current by a factor of 10, and, as mentioned above, changes in the dihedral angle of the two phenyl groups can also induce changes in the current. However, our results show that the SC-PEP molecules have significantly reduced blinking events than the SC$_{12}$ case in the measured frequency range. Therefore, we conclude that changes in the dihedral angle fall out of the measured frequency window.[63,64] We note that the dihedral angle can be chemically fixed to remove this degree of freedom and obtain stable conduction through the molecule.[61]

We note that others have performed noise spectroscopy of current time traces of single molecule junctions. [23,25,26,29,70–73] In those experiments, the $S(f)$ analysis was performed in the frequency range of 0.1–10$^6$ Hz, and the fluctuation of the tunneling current was attributed to changes in the molecule–electrode (Au-S) bond,[25] charge trapping at localized states (or defects),[70] or the change of molecule–electrode coupling through metal atom movements.[26,73] In contrast, we performed $S(f)$ analysis of the light emission from large-area STJs (in the frequency range of 10$^{-3}$ to 1 Hz) which provides a new way to visualize and examine the local current fluctuations within a large-area STJ, which is not possible with noise spectroscopy based on the tunnel current measured across the whole junction.

In our experiments, leakage radiation microscopy allows us to directly visualize the effective electrical contact of the STJs. This method, however, is diffraction limited and therefore, in principle, we cannot be certain whether a spot represents conduction via a single or a bundle of molecules. To deepen the understanding regarding the effective contact area, we evaluate the conductance of a single emitting spot. In the low bias region ($V < 0.3$ V), the $J(V)$ curves are linear from

![Figure 4. Representative emission intensity time traces of the STJs with a) SC$_{12}$ and b) SC-PEP SAMs recorded at an applied bias of −2.0 V at 0.5 s time intervals. The corresponding $S(f)$ analysis of the two emission intensity time traces is plotted in (c) and (d).](image-url)
which we determined a conductance of the SC-12 and SC-PEP junctions of $1.2 \times 10^{-7} G_0$ and $8.2 \times 10^{-7} G_0$, respectively ($G_0$ is the quantum conductance of $7.75 \times 10^{-5}$ S). Considering the number of emitting spots (the spot number at $-2.0$ V), that is $42 \pm 5$ for SC-12 and $8 \pm 3$ for SC-PEP junctions (the error bars represent the average determined from $>100$ measurements; see the Supporting Information), the conductance of each emitting spot is $2.9 \times 10^{-9}$ and $1.0 \times 10^{-7} G_0$, respectively. These conductance values are similar (within one order of magnitude) to previously reported values obtained from junctions ($10^{-9}–10^{-8} G_0$ for the SC-12 molecule\cite{33,74–77} and $10^{-7}–10^{-5} G_0$ for the SC-PEP molecule without the CH$_2$ unit between the sulfur and the phenyl unit\cite{76,78–82}) where the SAMs were contacted with a conducting atomic force microscope tip; in these junctions, the SAMs are chemisorbed on the bottom electrode and form a physisorbed contact with the top electrode. Thus, it seems that each spot represents conduction via a single, or very few, molecule(s), a finding that confirms our earlier suggestions\cite{20,28}.

Our results also reveal that the top electrode forms a 5.2 times more efficient contact with the SC-12 than with the SC-PEP SAMs in terms of effective contact area (i.e., the number of spots). We believe that this difference can be explained by the different stiffness of the molecules as molecules with a large conformational degree of freedom can form a better conformal contact with the top electrode than stiff molecules. For techniques that require spin coating of top-electrodes or protective barriers (that are placed between the SAM and the top electrode), such as organic semiconductors\cite{83} or graphene (oxide) inks\cite{84} from solution, differences in surface tension may also play a significant role in SAM//electrode interface formation.

3. Conclusions

In conclusion, we demonstrate control over the intermittent light emission of STJ-based plasmon sources by changing the flexibility of the molecular backbones. Stable light emission (originating from radiative decay of surface plasmons) is obtained for junctions with stiff conjugated molecules, in contrast to the same junctions but with aliphatic molecules. We found that light emission from STJs with aromatic molecules follows a $1/f$ behavior, while STJs with aliphatic molecules show an extra $1/f^2$ component representing the blinking induced by the conformational changes of the molecules. In addition, our methods allow for direct visualization and quantification of the effective electrical contact area making quantitative

Figure 5. The bias dependency of the intermittent light emission of junctions with SC-12 SAMs. a) Time traces of the light emission intensity at different biases. b) $S(f)$ analysis of the time traces in (a). c) The corresponding histograms of the $f_0$ with Gaussian fits to these histograms. The histograms are obtained from the measured emitting spots of four samples using videos consisting of 2000 frames.
comparisons to single molecule experiments possible and dynamic characteristics of the molecule–electrode interfaces of molecular tunnel junctions in operando. Thus, we disentangled two sources of heterogeneity—spatial and temporal—and showed that rigidity of the molecules—an inherent property of all molecules—within the junction directly influences the light emission during electron tunneling at the electrical contact areas.

We believe that our results are important for future studies where, for instance, strong intermolecular interactions could be explored to reduce blinking, or how molecular dipoles or charging of the SAMs via redox reactions could affect the molecule–electrode interactions and consequently the blinking dynamics.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

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