Dynamic response of glucagon/anti-glucagon pairs to pulling velocity and pH studied by atomic force microscopy

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Abstract

We used atomic force microscopy (AFM) to measure the unbinding force between antigen coupled to an AFM tip and antibody coated on the substrate surface. Dynamic responses of glucagon/anti-glucagon pairs with multiple pull-off steps to pH and pulling velocity were studied by AFM. Force–distance curves of a specific glucagon–anti-glucagon interaction system with mono-, di-, and multi-unbinding events were recorded, which may be attributed to a single, sequential or multiple breaking of interacting bond(s) between glucagon and anti-glucagon. We studied the dynamic response of glucagon–anti-glucagon pairs to various pulling velocities (16.7–166.7 nm/s). It was found that the mean value of the unbinding force was shifted toward higher values with increasing pulling velocity at each pH. This indicates that the friction force between glucagon and anti-glucagon may contribute to the unbinding force. Moreover, the dynamic response of glucagon–anti-glucagon pairs to pH (4–10) with different pulling velocities was studied. Within the acid range, the bond strength between the glucagon/anti-glucagon complex showed a rapid increase from pH 4 to 7 and reached a maximum (256.4 ± 48.9 pN at 166.7 nm/s) at neutrality, followed by a sharp decrease with increasing pH (pH 7–10). This could be attributed to the conformational change that occurred in glucagon when the pH value in solution was varied from the reference level at neutrality. This study demonstrated that the pH dependence of multiple antigen-antibody bond-rupture forces could be measured by a force-based AFM biosensor. Unraveling the relationship between inter-molecular force and intra-molecular conformational change in acid, neutral, and alkaline environments may provide new directions for future application of force measurements by AFM in proteomics or in the development of a clinical cantilever-based mechanical biosensor.

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1. Introduction

Atomic force microscopy (AFM), a nano-resolution technique that needs no complex preparation procedure for biological samples (Binnig et al., 1986), has proved to be a suitable approach to structural studies of single biomolecules (Sitko et al., 2003; Chen et al., 2004; Kuznetsov et al., 2005) and to the measurement of molecular forces at the single-molecule level. AFM was employed successfully for measurement of intra-molecular unfolding forces of individual proteins (Desmeules et al., 2002; Weisel et al., 2003) and inter-molecular forces between various ligand and receptor pairs (Meadows et al., 2003; Kaur et al., 2004; Kienberger et al., 2005). Compared to other biophysical approaches to direct measurements of inter-molecular forces, such as optical tweezers, the magnetic torsion device, and the biomembrane force probe (BFP), AFM has the advantage of high spatial resolution for detecting molecular interaction at high lateral resolution combined with immense force sensitivity in a physiologic environment.

For measurements of inter-molecular receptor/ligand binding strengths, ligands are attached to AFM tips and receptors are attached to substrate surfaces (or vice versa). The AFM tip is brought into contact with the surface so as to form receptor/ligand complexes. When the AFM tip is retracted from the
2.1. Functionalisation of AFM tip and substrate

AFM tips (CONTR™ Nanosensors, Germany) were coupled with glucagon (G3157, Sigma, CA) and a glass surface (Superfrost, Germany) coated with monoclonal anti-glucagon IgG (G2654, Sigma, CA). AFM tips were cleaned for 30 min in (Superfrost, Germany) coated with monoclonal anti-glucagon IgG (100 g/ml in PBS, pH 7.2) bound covalently to the glass substrate surface, the force required for dissociation (unbinding force), defined as the rupture force at the moment of sample separation, can subsequently be recorded and measured by pulling on the complexes until the interaction bond breaks. Such experiments allow for estimation of affinity, rate constants, and bond width of the binding pocket (Cai and Yang, 2003; Kokkoli et al., 2004; Kienberger et al., 2005), and even provide a tool for mapping the energy landscape (Cai and Yang, 2003; Lee and Marchant, 2003). In addition, these equipment provide insights into the molecular dynamics of the receptor/ligand recognition process by varying the loading rate of force application (Lee and Marchant, 2003; Kokkoli et al., 2004; Kienberger et al., 2005).

Glucagon is a single polypeptide which consists of one linear peptide with 29 amino acid sequences, and which exists at low concentrations as a monomer of low α-helix content (Borpmr et al., 1957; Gratzer et al., 1972). For understanding structure–function relationship, glucagon in solution was investigated by use of circular dichroism (Gratzer and Beaven, 1969; Gratzer et al., 1972) and 1H nuclear magnetic resonance (Boesch et al., 1978). The results obtained show that the concentration-dependent equilibrium of glucagon in mildly alkaline solution involves the formation of associated forms of higher α-helicity. In acid solution, the effect is still present, but is greatly diminished. These observations point out that the conformational properties of glucagon depend strongly on the pH of the solution conditions. Moreover, the biological function of glucagon is to maintain the blood sugar at a stable level. Quantitative observations are needed for determination of the interaction dynamics of glucagon with its antibody, anti-glucagon immunoglobulin G (IgG), for the purpose of developing a clinical glucagon sensor chip for predicting the risk of future cardiovascular events. Here, we report on the application of AFM to studies of the molecular dynamics of the glucagon/anti-glucagon recognition process for use in developing a clinical cantilever-based mechanical biosensor.

2. Materials and methods

2.1. Functionalisation of AFM tip and substrate

AFM tips (CONTR™ Nanosensors, Germany) were coupled with glucagon (G3157, Sigma, CA) and a glass surface (Superfrost, Germany) coated with monoclonal anti-glucagon IgG (G2654, Sigma, CA). AFM tips were cleaned for 30 min in H₂SO₄/H₂O₂ (70:30, v/v) and then extensively rinsed with distilled water. The tips were then silanised with a 5% solution of 3-aminopropyltriethoxysilane (Fluka Chemie, Switzerland) in 5% ethanol solution for 30 min at room temperature, followed by several rinses with a solution of 5% ethanol/95% distilled water. The tips were then immersed in a 2.5% glutaraldehyde solution in phosphate-buffered saline (PBS, pH 7.2) for 1 h and then extensively rinsed with PBS solution. In the last step, the glucagon solution (100 µg/ml in PBS, pH 7.2) bound covalently to the AFM tip via the amino groups after incubation overnight at 4 °C. Before being used in the acquisition of force measurements, glucagon-functionalised tips were rinsed with PBS. Anti-glucagon IgG (100 µg/ml) was covalently to the glass substrate by use of the method described by Vinckier et al. (1998). The surface coverage of sample IgG on the glass and glucagon on the tip were measured by use of an AnaLight® Bio200 dual polarization interferometric biosensor (Farfield Sensors Ltd., Salford, UK).

2.2. AFM force measurements and data analysis

All force measurements were performed with a SPA-300HV scanning probe microscope (SPM) controlled by an SPI 3700 probe station unit (Seiko Instruments Inc., Chiba, Japan). All experiments were carried out in a liquid cell filled with freshly prepared aqueous solution at room temperature. Commercial AFM tips (silicon CONTR™ Nanosensors, Germany) with a probe radius of 20 nm were modified as described above. Each cantilever was individually calibrated by a thermal fluctuation method (Lévy and Maaloum, 2002) for determining its spring constant. The average spring constant for the cantilevers employed here was 80 ± 10 pN/nm. Moreover, the sensitivity of an individual cantilever was calibrated by use of the slope of the cantilever deflection (mV) versus deflection distance (nm) curve when the cantilever was applied on a stiff surface. The sensitivity value was found to be in the range from 11.4 to 18.3 mV/nm.

During a force measurement cycle (see Fig. 1), the piezo-scanner is moved toward the AFM tip by piezo-expansion, and the velocity constant is kept until the scanner is brought into contact with the tip (point 2 in Fig. 1). As the forward motion continues, the cantilever is pressed into the sample surface until a point of maximum load is reached (point 3). The direction of motion is then reversed, and the piezo-scanner is withdrawn from the AFM tip. The deflection distance of cantilever bending was limited to less than 60 nm for all force measurements. During the retract portion of the force measurement, i.e. when the piezo-scanner is withdrawn from the tip surface, the tip adheres to the surface due to interactions between the tip and the sample. The strength of the unbinding force is calculated from the difference between the maximum cantilever deflection (point 4) during the retraction process of the curve and the point of zero cantilever deflection (point 1). In this work, we set the piezo moving displacement with 500 nm z-amplitude at moving time of 3, 5, 10, and 30 s, and thus obtaining the pulling velocity of 16.7, 50, 100, and 166.7 nm/s, respectively. To measure the unbinding force made as function of pulling velocity, the velocity of piezo-scanner was varied from 16.7 to 166.7 nm/s. More than 100 individual force measurements for each vertical velocity of piezo-scanner were recorded at each pH.

3. Results and discussion

To understand the dynamic response of individual glucagons/anti-glucagon pairs to pH, we used AFM to measure the unbinding force required to separate complementary glucagon/anti-glucagon pairs at different pH in aqueous solution. Force measurements of the glucagon–anti-glucagon interactions were carried out by use of an AFM in the force scan mode. For these experiments, a glass substrate, coated with
monoclonal anti-glucagon IgG, was brought in contact with an AFM tip coupled with glucagon (Fig. 1A). The unbinding force of the glucagon–anti-glucagon interaction was measured as the piezo-scanner was retracted from the AFM tip surface. A schematic diagram of a typical unbinding force measurement cycle between glucagon and anti-glucagon is shown in Fig. 1B. The maximal cantilever deflection, resulting from a sharp adhesive ‘pull-off’ event (point 4), during the retraction trace of the force measurement is related directly to the magnitude of the unbinding force required for breaking the glucagon–anti-glucagon bond(s) formed (Allen et al., 1999).

Fig. 2 shows representative examples of the force–distance curves acquired with these systems. No noticeable pull-off event was observed in the tip–glass system (Fig. 2A). In the glucagon–anti-glucagon system (Fig. 2B), however, a clear force discontinuity was present as a result of the adhesion force (unbinding force) between the glucagon–anti-glucagon pair. This suggests that the force measured in the glucagon–anti-glucagon system can be attributed to the specific glucagon–anti-glucagon unbinding event. Moreover, to confirm that glucagons–anti-glucagon binding caused the observed force discontinuity, the anti-glucagon surface was flooded with glucagon as a control experiment effectively to block the combining sites of the anti-glucagon IgG on the substrate. The recorded force curves displayed no adhesion pull-off event (or no discrete adhesion point), i.e. an unbinding force was not observed (see Fig. 2C). The lack of discrete adhesion points in the force curves obtained in the two control experiments (Fig. 2A and C) highlights the specificity of the interaction between the glucagon coupled to the tip and the anti-glucagon coated to the substrate.

In Fig. 2B, a slope change before the adhesion point appeared during the retraction process may be attributed to a decrease
Fig. 2. Measurements of unbinding forces. Force vs. distance curves of no adhesive force appeared in the retraction process, which were obtained by use of (A) tip/glass and (C) glucagons/glucagon-occupied anti-glucagon systems as two control experiments. (B) A typical force–distance curve of the glucagon–anti-glucagon system with a specific unbinding event that appeared in the retracting process. (D) Another type of recorded curve, which kept the same slope during the retraction process and was thought to be a non-specific interaction between the tip and the substrate.

The effective stiffness $k$ of the system can be expressed by $k = (k_{\text{cant}} \times k_{\text{inter}})/(k_{\text{cant}} + k_{\text{inter}})$, where $k_{\text{cant}}$ and $k_{\text{inter}}$ is the stiffness of the cantilever and glucagon/IgG complex, respectively. If $k_{\text{inter}}$ is high (specific interaction) as compared to $k_{\text{cant}}$ the effective stiffness is equal to the cantilever stiffness and then quasi constant which is the case in Fig. 2B. In the other case (non-specific interaction) the value of the effective stiffness is close to the interaction stiffness and then low and not constant, as shown in Fig. 2D. The non-specific force curve shown in Fig. 2D could be attributed to two factors. First, the tip and the substrate were not completely coupled with glucagon and anti-glucagon, respectively. When the tip was pressed onto the substrate, the linker (aminopropyltriethoxysilane + glutaldehyde) on the tip non-specifically contacted with the linker or anti-glucagon on the substrate (Willemsen et al., 1998). Second, the time of contact of the tip with the substrate was too short for the glucagon on the tip to have enough time to complete the binding process with anti-glucagon before the approach process became a retraction process (Wong et al., 1997; Willemsen et al., 1998). As a result, specific and non-specific interaction can be distinguished by the change in slope during the retraction process.

Fig. 3 shows the force–distance curves of the glucagon–anti-glucagon system with mono- (Fig. 3A), di- (Fig. 3B) and multi- (Fig. 3C) unbinding events (pull-off steps) recorded for a pH 7 solution, which may be attributed to a single, sequential, or multiple breaking of interacting bond(s) between glucagon and anti-glucagon (Allen et al., 1997; Lo et al., 1999). In this study, an AFM tip with a probe radius of about 20 nm was coupled with glucagon and glass surface coated with anti-glucagon IgG, which has been shown to be a Y-shaped molecule of overall dimensions approximately $14.2 \times 8.5 \times 4.0$ nm by X-ray crystallography studies (Ahluwalia et al., 1992). In addition, a surface coverage of IgG of $9.47 \times 10^{-15}$ mol/mm$^2$ (or an area per IgG molecule of $158.4$ nm$^2$/molecule), the most likely surface structure is that of a monolayer. Such preparations of a monolayer IgG and glucagon surface were then employed in the force measurements of specific glucagons–anti-glucagon interactions (Fig. 3A–C). Schematic representations of the A, B, and C binding events are shown in a, b, and c, respectively. The force curve with one adhesion point (Fig. 3A) indicates that the single-pair interaction was broken in the retraction process of the force measurement. Such a force curve could be attributed to the fact that the anti-glucagon IgG molecules randomly immobilized on the glass could be oriented (a1) or lying on the substrate (a2) with one binding site.
Fig. 3. Force–distance curves of the glucagon–anti-glucagon system with (A) mono-, (B) di-, and (C) multi-unbinding events (pull-off steps) recorded in pH 7 solution at a pulling velocity of 166.7 nm/s, which may be attributed to (A) a single, (B) sequential, or (C) multiple breaking of bond(s) between the tip and the substrate. Schematic representations of the A, B, and C binding events are shown in a, b, and c, respectively. The anti-glucagon IgG molecules randomly immobilized on the glass could be oriented (a1) or lying on the substrate (a2) with one binding site oriented to bind a single glucagon on the tip. Two separate glucagons on the tip can bind to two identical combining sites on one anti-glucagon (b1) or to separate combining sites on two anti-glucagon molecules (b2). Several separate glucagons on the tip can bind to three (c1) to four (c2) separate combining sites on anti-glucagons.

In Fig. 3B, the force curve with two adhesion points indicates a sequential breaking of interacting bond(s) between the tip and the substrate. Two separate glucagons on the tip might bind to two identical binding sites on one anti-glucagon (b1) or to separate binding sites on two anti-glucagon molecules (b2) when the tip is in contact with the substrate surface. In addition, in some force curves (Fig. 3C), there were multiple pull-off steps in the retraction process that could be attributed to multiple breaking of interacting bonds between the tip and the substrate. The multiple pull-off steps observed could be explained by two factors. First, glucagon is a small molecule (MW = 3485), and more than three separate glucagons on the tip might bind to three (c1) to four (c2) separate binding sites on anti-glucagons when the tip is in contact with the substrate surface. Second, such multiple pull-off steps that appeared in a retracting process may be due to partial unbinding forces related to conformational transitions in the binding sites of anti-glucagon and induced protein-unfolding events (Lee and Marchant, 2003). As discussed above, the multiple unbinding events that were measured could result from the non-specific interaction or protein unfolding, therefore, they were neglected in our calculations.

To investigate the dependence of the pulling velocity of glucagon–anti-glucagon unbinding forces, we recorded and plotted all specific unbinding events as force histograms, which were then subsequently subjected to Gaussian fitting for analysis (Evans and Ritchie, 1997; Lee and Marchant, 2003; Bartels et al., 2003; Heymann and Grubmüller, 1999). Fig. 4 shows histograms of the unbinding force between a glucagon tip and an anti-glucagon substrate measured with various pulling velocities (V) of 16.7, 50.0, 100.0, and 166.7 nm/s, respectively, at pH 7. For each unbinding force (F) obtained, more than 100 unbinding events from about 500 approach/retraction cycles were recorded. All histograms were fitted to a Gaussian function (Evans and Ritchie, 1997). The mean value of the Gaussian equals the most probable unbinding force, with statistical errors given by standard deviation. All AFM force measurements were carried out in PBS at 25 °C.

Fig. 4. Histograms of the unbinding force between a glucagon tip and an anti-glucagon substrate measured with various pulling velocities (V) (16.7, 50.0, 100.0, and 166.7 nm/s) in pH 7 solution. For each unbinding force (F) obtained, more than 100 unbinding events from about 500 approach/retraction cycles were recorded. All histograms were fitted to a Gaussian function (Evans and Ritchie, 1997). The mean value of the Gaussian equals the most probable unbinding force, with statistical errors given by standard deviation. All AFM force measurements were carried out in PBS at 25 °C.
solution. For each unbinding force ($F$) obtained, more than 100 unbinding events from about 500 approach/retraction cycles were recorded. The mean unbinding force of the glucagon–anti-glucagon complex at a given pulling velocity was determined from a force histogram obtained from more than 100 measurements. The corresponding mean unbinding forces, obtained by fitting the histogram to a Gaussian function, were 151.0 ± 22.0, 170.6 ± 26.1, 210.4 ± 38.0, and 256.4 ± 48.9 pN at pulling velocities of 16.7, 50.0, 100.0, and 166.7 nm/s, respectively.

In Fig. 4, the mean value of the unbinding force ($F$) was shifted toward higher values with increasing pulling velocity at pH 7. The fact that the unbinding force increased with increased pulling velocity indicates that the friction force between glucagon and anti-glucagon may contribute to the unbinding force (Heymann and Grubmüller, 1999). We assume the difference between the unbinding force obtained at each pulling velocity and the unbinding force at a zero pulling velocity corresponds to the friction effect. The assumption is supported by a linear relationship between the unbinding force and the pulling velocity (Heymann and Grubmüller, 1999). This shows that the unbinding force is proportional to the pulling velocity ($F = \alpha V$, where $\alpha$ is the friction coefficient and $V$ is the pulling velocity) (Heymann and Grubmüller, 1999).

The friction coefficient with varied pulling velocity at each pH can be calculated from the slope of each plot. The friction coefficient obtained was 0.1598, 0.4239, 0.7137, 0.3596 and 0.2217 pN/s/nm at pH 4, 6, 7, 8, and 10, respectively. The maximum friction coefficient was obtained at pH 7, and all of the friction coefficients decreased as the pH value both increased and decreased. This phenomenon has also been interpreted by Bell (1978) and Evans et al. (2001) based on diffusion theory. In addition, for force measurements in a liquid environment, hydrodynamic drag may make a considerable contribution to the force that is probed, especially at high pulling velocity (Lee and Marchant, 2003). To understand the possible effect of hydrodynamic drag on the unbinding force, we measured the deflections of the free-standing AFM cantilever as a function of distance in PBS buffer and recorded them in the non-contact region at various pulling velocities in the range of from 16.7 to 166.7 nm/s. Within this range of pulling velocities, it was found that the signals of the reflected laser beam (deflections) in the non-contact region in the force measurement cycle were kept constant, with slopes close to zero in a plot of deflection versus position (data not shown). This indicated that hydrodynamic drag did not have any influence on the unbinding force in the force measurements at various pulling velocities in this work.

Fig. 6 summarizes the dynamic response of glucagon–anti-glucagon pairs to the pH value (4–10) with different pulling velocities (16.7–166.7 nm/s). Within the acid range (pH 4–7), the unbinding force of the complex showed a rapid increase with increasing pH, reaching a maximum (256.4 ± 48.9 pN at 166.7 nm/s) at neutrality (pH 7), followed by a sharp decrease with increasing pH (pH 7–10). This indicates that the bond strength between glucagon and anti-glucagon in acidic or alkaline solution is reduced compared to that in a neutral solution. These results could be attributed to two factors below. First, a conformational change may occur in glucagon when the pH value in solution is changed from the reference level at neutrality. The glucagon in aqueous solution does not have a stable α-helical structure, which is essentially dependent on the pH over the range in which glucagon is soluble. The concentration-dependent equilibrium of glucagon in mildly alkaline solution involves the formation of associated forms of higher α-helicity. In acid solution, the effect is still present, but, is greatly diminished (Gratzer and Beaven, 1969; Gratzer et al., 1972; Panijpand and Gratzer, 1974). These observations point out that the conformational properties of glucagon depend strongly on the pH of the solution. This may reduce the binding strength between glucagon and anti-glucagon, and thus it led to a decrease in the

Fig. 5. Pulling-velocity dependence of the unbinding force for glucagons–anti-glucagon pairs. It clearly shows that the unbinding force is proportional to the pulling velocity at various pH values (4, 6, 7, 8, 10) ($R^2 = 0.9585, 0.9911, 0.9982, 0.9440$ and $0.9859$ for pH 4, 6, 7, 8 and 10, respectively).

Fig. 6. pH dependence of the unbinding force for glucagons–anti-glucagon pairs with different pulling velocities (16.7–166.7 nm/s). The error bars correspond to one standard deviation, as indicated in Fig. 4, and are placed on all data points. Within the acid range (pH 4–7), the unbinding force of the complex showed a rapid increase with increasing pH and reached a maximum (256.4 ± 48.9 pN at 166.7 nm/s) at neutrality (pH 7), followed by a sharp decrease with increasing pH (pH 7–10).
unbinding force in both the acid and alkaline range. Second, the biological function of glucagon varies with the pH value. When the force of glucagon–anti-glucagon pairs was measured in alkaline solution, the glucagon rapidly converted to monode-samidoglucagon (Gratzer et al., 1972). This result could lead to a decrease in the biological binding strength of glucagon with anti-glucagon. On the other hand, when the force measurement was performed in acid solution, the glucagon aggregated to produce gels (Panijpan and Gratzer, 1974), this also could weaken the interaction force between glucagon and anti-glucagon.

4. Conclusions

We have shown the advantage of using AFM, which is a sensitive force probe for elucidating the molecular details of specific unbinding events between glucagon and anti-glucagon with different pulling-velocities at various pH values. A specific glucagon–anti-glucagon interaction system with mono-, di-, and multi-unbinding events was found in the force–distance curves, which may be attributed to a single, sequential, or multiple breaking of interacting bond(s) between glucagon and anti-glucagon. Moreover, each average unbinding force of glucagon–anti-glucagon pairs was shifted toward higher values with increasing pulling velocity (16.7–166.7 nm/s) at various pH values (4, 6, 7, 8, 10). This finding indicates that the friction force between glucagon and anti-glucagon may contribute to the unbinding force. Furthermore, the dynamic response of glucagon–anti-glucagon pairs to pH levels (between 4 and 10) was studied. We found that the bond strength between the glucagons/anti-glucagon complex showed a rapid increase from pH 4 to 7, and reached a peak at neutrality, followed by a sharp decrease with increasing pH (pH 7–10), which could be attributed to the conformational change that occurs in glucagon when the pH value in solution is varied from the reference level at neutrality. Our results demonstrate that the pH dependence of multiple antigen-antibody bond-rupture forces could be correlated with intra-molecular conformational change in acid, neutral, and alkaline environments.

References


