Acquiring Biological Information of Individuals Using Quantum Mechanics

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Abstract
We developed a single-molecule technology that measures the tunneling currents induced by single-base molecules of DNA, RNA, and peptides passing between two nanoelectrodes. Unlike existing sequencing methods, our technique can potentially sequence chemically modified bases and amino acids, many of which are important disease markers.

1. Introduction
Many people expected that the end of the Human Genome Project would mark the starting point of personalized medicine and therapeutics based on genomic information. However, the cost and throughput to read out a human genome have been a substantial barrier to the realization of this technology. Our single-molecule electrical sequencing technology is based on semiconductor technologies and is expected to become an ultimate sequencer with ultra-low cost and ultra-high speed. Unlike existing sequencing technologies, single-molecule sequencing technology is expected to identify sequences of base molecules in RNA and sequences of amino acid molecules in peptides because it can identify small differences in molecular electronic structures through differences in single-molecule conductances. In addition, single-molecule sequencing technology may be able to detect chemically modified base molecules and amino acid molecules that are known markers of diseases such as cancer; there are no existing technologies detecting the chemical species directly. We hope to identify base molecules and chemically modified base molecules in DNA and RNA as well as amino acid molecules with and without chemical modifications using single-molecule sequencing technology.

2. Methods
All single-molecule electrical measurements were performed using nanofabricated mechanically controllable break junction (MCBJ) electrodes, in which a metal wire produced on an insulating substrate is fractured by pushing a piezo actuator up, forming gaps of less than several nanometers [1]. The nanoelectrode spacing was configured to be 0.8 nm for DNA and RNA and 0.5 nm and 0.7 nm for peptides because, unlike base molecules of DNA and RNA, the molecular sizes of amino acid molecules differ. Current–time profiles were measured at a sampling rate of 10 kHz under an applied potential of 0.4–0.7 V when a solution containing analytes was dropped onto the nanogap electrodes. Sequences of base molecules and amino acid molecules were determined on the basis of current–time profiles using in-house analysis algorithms.

3. Results and Discussion
We first determined the single-molecule conductances of base molecules of DNA and RNA. When current–time profiles were measured, spike-like signals were observed and were characterized on the basis of their maximum currents and current duration. We explored the electron transport mechanism through single-base molecules to determine the voltage dependence of the electrical currents. We observed that the electrical currents increased linearly with increasing applied voltage, indicating that the electrical currents originated from tunneling currents between nanoelectrodes. The single-molecule conductance histograms showed single peaks, and the single-molecule conductance order in DNA was guanine (G) > adenine (A) > cytosine (C) > thymine (T), whereas that in RNA was G > A > C > U [2]. This result indicates that electric currents allow us to identify single-base molecules.

Because single-molecule conductances strongly depend on the highest occupied molecular orbital (HOMO) energies, single molecules with different HOMO energies can be distinguished even though the differences in HOMO energies are small. Although methylated cytosine (mC) is a well-known disease marker, no existing sequencing technologies can identify it directly. The single-molecule conductance of mC is larger than that of C because of a small difference of 0.1 V in their HOMO energies [3]. In addition...
Fig. 2 Sequences of base molecules in DNA and RNA and sequences of amino acid molecules in peptides can be identified on the basis of single-molecule conductances of base and amino acid molecules, where tunneling currents between nanoelectrodes are measured.

to mC, oxo-G, which is a known cancer marker, can also be identified on the basis of single-molecule conductance. Therefore, our single-molecule identification method can detect chemically modified base molecules.

Next, we applied our single-molecule identification method to short DNA molecules such as TGT [4]. When the current–time profiles were measured, spike-like signals were observed in addition to the signals of single-base molecules. Two single-molecule conductance peaks corresponding to G and T were observed in the single-molecule conductance histograms. In the case of TGT, two lower-conductance plateaus were observed in the first and third lower positions, and the plateau associated with guanine was observed in the middle higher plateau. Therefore, we achieved single-molecule electrical sequencing based on single-molecule conductances. However, not only TGT but also G, T, and TG were observed because of stochastic traps caused by Brownian motions.

Using stochastic traps, we can expect to achieve a random sequencing method that is applicable to existing DNA sequencers. We attempted to perform single-molecule random sequencing of microRNA molecules, which are formed by 22–25 base molecules and are well known as cancer markers. As expected, different randomly fragmented sequences were obtained and assembled into sequence contigs, and sequence contigs were assembled into continuous sequences. Therefore, we successfully identified sequences of microRNA, which cannot be sequenced directly by existing DNA sequencing technologies.

Consequently, 12 amino acid molecules and phosphorylated tyrosine were identified on the basis of single-molecule conductances [5]. In addition to DNA and microRNA signals, spike-like signals were also observed; the single-molecule conductance histograms corresponding to the different amino acid molecules showed easily distinguishable peaks. Consequently, amino acid molecules of peptides can be identified using single-molecule electrical sequencing technology. The results of our experiments make us hopeful of using this platform to develop a high-throughput, ultra-fast, and inexpensive system for driving DNA sequencing beyond the limitations of current technologies.

4. Conclusions

Single-base molecules of DNA and RNA and single amino acid molecules of peptides can be identified using tunneling currents between nanoelectrodes. Single-molecule sequencing technology can determine sequences of base molecules in DNA and RNA and partial sequences of amino acid molecules in peptides. In addition, chemically modified base molecules and amino acid molecules can be distinguished via single-molecule conductances. The results of our experiments make us hopeful of using this platform to develop a high-throughput, ultra-fast, and inexpensive system for driving DNA sequencing beyond the limitations of current technologies.

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