In situ oligonucleotide synthesis on carbon materials: stable substrates for microarray fabrication

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Received August 21, 2007; Revised and Accepted November 27, 2007

ABSTRACT

Glass has become the standard substrate for the preparation of DNA arrays. Typically, glass is modified using silane chemistries to provide an appropriate functional group for nucleic acid synthesis or oligonucleotide immobilization. We have found substantial issues with the stability of these surfaces as manifested in the unwanted release of oligomers from the surface when incubated in aqueous buffers at moderate temperatures. To address this issue, we have explored the use of carbon-based substrates. Here, we demonstrate in situ synthesis of oligonucleotide probes on carbon-based substrates using light-directed photolithographic phosphoramidite chemistry and evaluate the stabilities of the resultant DNA arrays compared to those fabricated on silanized glass slides. DNA arrays on carbon-based substrates are substantially more stable than arrays prepared on glass. This superior stability enables the use of high-density DNA arrays for applications involving high temperatures, basic conditions, or where serial hybridization and dehybridization is desired.

INTRODUCTION

DNA arrays have become a vital component in genomic research for high-throughput gene expression analysis (1,2), mutation detection (3,4), gene discovery and genetic mapping studies (5) and protein–DNA interaction analysis (6–9).

Several methods for fabricating DNA arrays have been described (10). These fall into two groups: in situ synthesis (11–16), or deposition and immobilization of pre-synthesized DNA sequences (17–26). Immobilization of pre-synthesized oligonucleotides offers the flexibility needed to quickly make low and medium-density arrays containing anywhere from several dozen to several hundred features per array. Although the technologies and skills needed to prepare low density arrays are readily available to a large number of researchers, there is an increasing need for high-throughput array-based analyses with high density arrays (>50 000 features per square centimeter). Both photolithographic and ink-jet array synthesis methods prepare arrays in a combinatorial manner, one nucleotide at a time. This permits the end user to create a high-density array with a greater diversity of sequences in less time and at a lower cost than spotting pre-synthesized oligonucleotides.

Glass, with its low intrinsic fluorescence, non-porosity and ease of modification using silane chemistries has become the standard surface for fabricating DNA arrays (23). A disadvantage of glass substrates, however, is the intrinsic hydrolytic instability of the siloxyl linkage employed in glass chemical modification (27,28). Glass substrates are limited to mild pH conditions and moderate-to-low temperatures, limiting the applications of arrays fabricated on this material. While this may not affect common applications such as gene expression or SNP detection, which are formulated to work with current glass-based platforms, new substrates exhibiting greater stabilities under a broader range of conditions would enable new applications such as those employing extended high temperature incubations or harsh chemical conditions.

Several non-glass substrates including silicon (29–33), gold (34,35) and polymeric materials (36,37), have been reported for deposition of pre-synthesized oligonucleotides. While these substrates can offer added benefits such as increased conductivity or flexibility over their glass counterparts, they have yet to become a widely used alternative.

The use of carbon-based substrates for the fabrication of low-density hand-spotted DNA arrays (38–40) has also been described. DNA arrays fabricated on such carbon-based surfaces are extremely robust due to both the intrinsic chemical stability of the substrate and to the carbon–carbon bonds employed for surface attachment; arrays prepared on carbon-based substrates exhibit
greater stability than arrays prepared on either silicon or gold substrates (38,40).

In this study, we show the utility of carbon substrates for the in situ light-directed synthesis of DNA arrays. The stability of the resulting DNA arrays is dramatically increased compared to DNA arrays prepared on glass. This increased stability expands the utility of high-density arrays by enabling their use under higher temperature conditions and with extended reaction times and greater extremes of pH, as well as permitting their regeneration and reuse.

MATERIALS AND METHODS

All reagents were purchased from Sigma Aldrich and used without further purification unless otherwise stated.

**Silanization of glass slides**

A 0.1% acetic acid in 95% ethanol stock solution was prepared. ArrayIt SMC Superclean glass slides (Telechem International, Inc., Sunnyvale, CA, USA) were stirred in 2% (v/v) N-(3-triethoxysilylpropyl)-4-hydroxy-butyramide (Gelest, Inc., Morrisville, PA, USA) in stock solution for 4 h at room temperature. The slides were then rinsed by stirring in fresh stock solution for 15 min. After being rinsed three times in diethyl ether, slides were transferred to a pre-heated (120°C) oven for a minimum of 2 h, after which time they were cured under vacuum overnight. Slides were stored desiccated until ready for use (Scheme 1a).

**Glassy carbon material**

Custom sized (2.5 cm × 4.0 cm) glassy carbon plates (Sigradur K) were purchased from Hochtemperatur-Werkstoffe GmbH, Thierhaupten, Germany. Glassy (vitreous) carbon is a hard, ceramic-like material produced by the pyrolysis of carbon-rich polymers. The resulting material is composed of graphitic ribbons of pure carbon with fullerene-like microstructures (41,42). Like graphitic carbon, it is primarily composed of sp² carbons, although the density of glassy carbon is greater than that of its graphitic counterpart. Detailed information on this material can be found in the cited references as well as in several books on the subject of carbon materials (43,44).

**Preparation of nanocrystalline diamond thin films**

Nanocrystalline diamond samples were a gracious gift from Dr James Butler at Naval Research Laboratories. Thin films of 0.50–0.59 μm CVD diamond were deposited on n-type Si <100> at 850°C using 6.4 Torr methane at 2.50 Hz, 1000 W (45).

![Scheme 1](image-url)
Hydrogen termination of glassy carbon and diamond substrates

Glassy carbon plates and diamond substrates were chemically cleaned prior to hydrogen termination. Each substrate was sonicated for 5 min in CHCl₃ and cleaned with a series of acid treatments at 60°C for a minimum of 1 h first in 4:3:1 water:nitric acid:hydrochloric acid and then in 3:2 sulfuric acid:nitric acid. (Caution: these acid washes and associated fumes are highly caustic, proper protective equipment and appropriate ventilation must be employed.) In between acid treatments, the substrates were rinsed with deionized (DI) H₂O. Prior to functionalization, glassy carbon and diamond surfaces were heated to ~900°C under vacuum and treated with a 13.56 MHz inductively coupled H-plasma for 20 min at 50 Torr to generate a hydrogen terminated surface (40) (Scheme 1b).

Generation of free alcohol groups on carbon substrates

Following hydrogen termination, the carbon surfaces were photochemically functionalized by placing 30 μl of 9-decene-1-ol directly onto the surface and covering with a clean quartz coverslip. The surfaces were irradiated under N₂ purge with a low-pressure mercury vapor quartz grid lamp (λ = 254 nm) for 8–12 h. After the photo-reaction, the surfaces were briefly rinsed with ethanol, DI H₂O and sonicated 2 × 5 min in CHCl₃ before being immersed in concentrated ammonium hydroxide for 2 min. The surfaces are then stored desiccated until ready for use (Scheme 1b).

In situ oligonucleotide synthesis

Light-directed photolithographic synthesis was performed with a digital micromirror-based Biological Exposure and Synthesis System (BESS) connected to a Perceptive Biosystems Expedite Nucleic Acid Synthesis System (Framingham, MA, USA) as described previously (12,16,46). Oligonucleotide synthesis was carried out using a modified DNA synthesis procedure, where the removal of the photolabile NPPOC (3’-nitrophenylpropyloxycarbonyl)-protecting group was achieved by irradiation with 3.95 J/cm² of 365 nm light from a 200 W Hg/Xe arc lamp (Newport, Stratford, CT, USA). The optimum dose for removal of the NPPOC protecting group was determined empirically (data not shown). The UV-irradiation time was determined by measuring the lamp power (in mW/cm²) at 365 nm and adjusting the exposure time to ensure a dose of 3.95 J/cm².

Oligonucleotide synthesis reagents [DCI activator, acetonitrile (dry wash and amidite diluent) and oxidizer solution] were purchased from Sigma-Proli go; exposure solvent was purchased from Nimblegen Systems Inc. (Madison, WI, USA). All anhydrous reagents were kept over molecular sieves (Trap Packs, Aldrich).

All NPPOC-protected phosphoramidites [5’-NPPOC-dAdenosine(tac) 3’-β-cyanoethylphosphoramidite (dA), 5’-NPPOC-dThymidine 3’-β-Cyanoethylphosphoramidite (dT), 5’-NPPOC-dCytidine(ib) 3’-β-cyanoethylphosphoramidite (dC), 5’-NPPOC-dGuanosine(ipac) 3’-β-cyanoethylphosphoramidite (dG)] were manufactured by Proligo Biochemie GmbH (Hamburg, Germany) and purchased from Nimblegen Systems Inc.; NPPOC-phosphoramidites were diluted (1 g in 60 mL) with dry acetonitrile (amidite diluent).

Addition of a NPPOC-protected phosphoramidite proceeds as follows: (i) after condensation of the previous NPPOC-protected base to the growing DNA strand, the synthesis flow cell (volume ~100 μl) is flushed with 500 μl of exposure solvent; (ii) a digital image (mask) representing the locations for the next base addition illuminates the surface with 3.95 J/cm². During irradiation of the array, the exposure solvent is constantly flowed through the flow cell at a rate of 100 μl/0.5 J/cm² to maintain sufficiently basic conditions (47) to drive the photo-catalysed elimination reaction. Following irradiation, (iii) the array is washed with acetonitrile (~400 μl) to remove residual exposure solvent, dry wash (~500 μl) to remove trace water, and activator solution (~100 μl). Coupling of the next base is achieved by filling the flow cell with a 1:1 solution of the desired phosphoramidite and activator. All 5’-NPPOC-protected amidites undergo a single 40s coupling step, whereas the Cy3 dye phosphoramidite [0.03 M, Glen Research, Sterling, VA, USA] is subjected to two 300s coupling steps. After amidite coupling, the array is washed with acetonitrile (~100 μl) and either oxidized by flushing the cell with oxidizer solution (THF, pyridine, iodine, water; ~500 μl) or subjected to the next phosphoramidite addition. The non-acidic conditions of deprotection (47) allow for oxidation of the backbone phosphate groups only after every fourth coupling step and at the end of the synthesis, rather than at every coupling step. We have found that this modification does not negatively impact sequence quality (Richmond, K.E., Rodesch, M.J., Kaysen, J. and Cerrina, F., unpublished data) and reduces both the synthesis time and the amount of waste generated.

After synthesis is completed, the nucleoside bases are deprotected in 1:1 ethylene diamine: absolute ethanol solution at room temperature for 2–4 h.

Table 1 contains the probe sequences synthesized on all low density arrays. The 3’ end of sequences 2–4 were separated from the surface by a 10-thymidine spacer, providing a distance of ~30 Å from the surface; this has been shown to increase hybridization efficiency (26,48). Probe 1 was terminally labeled with Cy3, while sequences complementary to probes 2–4 in Table 1 were modified at the 3’ end with a fluorescein label. For hybridization density determination, sequence 4 was used; all other data presented is only for sequence 1, in order to simplify the analysis.

<table>
<thead>
<tr>
<th>Probe sequence</th>
<th>Sequence: 3’ → 5’</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>TTTTTTTTTT-Cy3</td>
</tr>
<tr>
<td>2</td>
<td>(T₉)₃TTATTTGAAAACGTTGTCACC</td>
</tr>
<tr>
<td>3</td>
<td>(T₉)₃GTTATGAAAACGTTGTCACT</td>
</tr>
<tr>
<td>4</td>
<td>(T₉)₃GGCTACTGGAACGTCTCA</td>
</tr>
</tbody>
</table>
DNA hybridization and washing
Complementary oligonucleotides for probes 2–4 were purchased from IDT (Coralville, IA, USA) and the University of Wisconsin – Madison Biotechnology Center (Madison, WI, USA). All arrays were hybridized by placing 30 µl of the fluorescantly-tagged complement (1 µM, 1 x SSPE [10 mM NaH₂PO₄, 0.15 M NaCl, 1 mM EDTA, pH = 7.4], 45°C) on the surface, covering with a coverslip, and incubating for 1 hr in a humid chamber. Nonspecifically bound DNA was removed by incubating the surface in 1 x SSPE for 15 min at 37°C. Dehybridization was achieved by incubating the surfaces in 8 M urea (RT, 20 min). Unless otherwise noted, fluorescence scans were taken on a Genomic Solutions GeneTac UC 4 x 4 scanner (Ann Arbor, MI, USA). Arrays hybridized with fluorescein-labeled complementary DNA strands were scanned in 1 x SSPE.

Determination of hybridization density
Hybridization density was determined using a wash-off method as described previously (49). Arrays consisting of a single sequence (Sequence 4, Table 1) were synthesized over the entire (1 cm x 1.3 cm) synthesis area. After hybridization and washing, the array was transferred to 20 ml of wash-off buffer (40 mM KCl/132 mM KOH) in a 50 ml falcon tube and shaken vigorously for 15–20 min. Calibration solutions containing fluorescein-labeled target DNA (1 x 10⁻¹¹ to 1 x 10⁻⁸ M) were prepared in 42 mM KCl, 132 mM KOH. Using a fluorescence plate reader (BIOTEK, Flx 800, 200 µl per well), the fluorescence from the calibration solutions and unknown samples were measured and the density of hybridized DNA was calculated.

Thermal stability determination
In a stirring isothermal water bath, 40 ml of 2xSSPE with 0.2% (v/v) SDS was pre-warmed to either 45°C or 60°C in 50 ml falcon tubes fitted with stir vanes. After an initial scan, arrays were incubated at one of the indicated temperatures. At pre-determined time points (0, 0.5, 1–4 h in 1 h intervals, and 4–24 h in 2 h intervals) the arrays were removed from the solution, rinsed with RT 1xSSPE and hybridized before being scanned and returned to the warm 2 x SSPE–SDS solution to continue incubating.

Hybridization stability determination
Arrays were subjected to 20 hybridization cycles: hybridized as described earlier, scanned, incubated in 8 M urea at RT for 20 min, rinsed with water, 1xSSPE and another hybridization step was performed. Complete dehybridization was verified by fluorescence imaging.

Base stability determination
Arrays were incubated in a solution of NH₄OH (15%, 55°C). (Caution: produces noxious fumes, ensure appropriate ventilation and open carefully as containers pressurize under elevated temperatures.) Fluorescence scans of the arrays were taken at pre-determined time points (0, 0.5, 1–12 h) by rinsing the array with DI H₂O, 1xSSPE, and hybridizing as described earlier.

PCR stability determination
A master mix consisting of 0.05 U/ml PicoMaxx enzyme, 0.1 mM each dNTP, in 1xPicoMaxx buffer (all Stratagene), augmented with 0.1% BSA (Promega) and 0.1% Tween-20 (Pierce) was made. Arrays were hybridized and fluorescence images captured before being fitted with a hybridization chamber (Grace Biolabs) and 250 µl of pre-warmed solution. The arrays were placed in a thermocycler (MJ Research) and subjected to a PCR protocol that has been previously reported (50). Briefly, the array was initially heated to 94°C for 9 min followed by 50 cycles of 94°C for 45 s, 65°C for 3 min and 72°C for 4 min. Following this treatment, the arrays were rinsed with water, hybridized and scanned. After scanning, the process was repeated.

Comparison of array image quality
A high-density array using a one-in-four design was employed to evaluate high-density image quality. This consists of a single pixel-sized feature (16 µm x 16 µm) containing single probe species separated by a single pixel on all sides (one-in-four). Probe sequence (3’TGTGTTTCTGGTCCCACCAAGTACTACTACTG) was synthesized as described earlier with UV-deprotection dose varying from 0.7 to 12.1 J/cm². Following synthesis and nucleic acid deprotection, the array was hybridized with a Cy3 tagged complementary DNA target for 1 h and imaged using a Nikon E800 fluorescent microscope fitted with a cooled CCD imaging system and imaging software by Metamorph (Universal Imaging Corporation).

RESULTS
Two-color fluorescence images of identical array designs on three different substrates are shown in Figure 1. Sequence 1 is terminally labeled with Cy3 and is presented in false-color red, sequences 2–4 are hybridized with their respective fluorescein-labeled perfect match complementary sequence (false-color green). While all three substrates exhibit similar Cy3 intensities and comparably low background fluorescence, all sequences hybridized on glassy carbon (b) exhibit a greater fluorescence than those hybridized on either glass or diamond. This is likely an artifact due to optical effects from the scanner used to image the array—in a similar comparison (see Array image quality section) using a fluorescence microscope, diamond showed greater fluorescence while glass and glassy carbon were comparable. This is consistent with the hybridization density results given subsequently.

Hybridization density
The hybridization density of a surface is a measure of the number of probe oligomers that are accessible to bind complementary DNA. Wash-off studies in which DNA is first hybridized to and then eluted from the surface for measurement in solution are a preferred method of
analysis as substrate-specific and any quenching effects are minimized or eliminated. The hybridization densities determined for all three surfaces are shown in Table 2. The density of fluorescently labeled complementary oligonucleotides hybridized to the surface was between 2 \times 10^{12} (4 pmol/cm^2) and 4 \times 10^{12} molecules/cm^2 (7 pmol/cm^2). These are 12–52% lower than densities reported for oligonucleotides immobilized on gold (5 \times 10^{12} oligonucleotides/cm^2, 8 pmol/cm^2) (49) and are 14–26% below the theoretical maximum oligonucleotide density of 1.7 \times 10^{13} oligonucleotides/cm^2 (17 pmol/cm^2) as calculated when the dsDNA helices are assumed to be tightly packed cylinders with diameters of 2 nm each.

### Thermal stability

To increase the specificity of hybridization it is often desirable to incubate arrays at higher temperatures. Such temperatures increase the rate of hydrolysis and lead to problems with array stability. To investigate this issue, the surface stability of arrays prepared on glassy carbon, diamond and silanized glass was compared at both 45°C and 60°C (Figure 2).

Under all conditions, glassy carbon and diamond exhibit greater stability than glass. After 24 h of incubation at either 60°C or 45°C, glassy carbon retains 99 ± 1% of the initial fluorescence signal. Arrays prepared on diamond retain 80 ± 5% of the initial fluorescence signal at 45°C and 67 ± 5% at 60°C after 24 h of incubation. Following 20 h of incubation at 45°C glass retains <5% of initial fluorescence and a similar loss is observed after only 6 h at 60°C.

### Base stability

A similar trend is seen when the substrates are immersed in 15% ammonium hydroxide at 55°C. After 12 h of incubation, the glassy carbon arrays retain 100 ± 3%
of their initial signal and diamond-based arrays retain 78 ± 4% of the initial signal. Fluorescence signals from arrays on glass substrates have < 5% of their initial signal after 1 h and exhibit no detectable signal after 4 h of incubation (Figure 3).

**Stabilities to multiple hybridization cycles**

Incubation in solutions containing either high salt concentrations or helix destabilizing reagents are standard methods to achieve DNA dehybridization. Incubation in 8M urea for 20 min at room temperature is relatively mild, yet sufficient to completely dehybridize probe and target molecules (data not shown). The changes in fluorescence signal for multiple hybridization cycles separated by incubations in 8M urea solution for 20 min were monitored (Figure 4). After 20 cycles, glass exhibited 31 ± 1% and glassy carbon exhibited 91 ± 5% of the initial fluorescence, while diamond exhibited no measurable loss of fluorescence (108 ± 7%).

**Stability to PCR cycles**

Glass showed very poor stability when subjected to the conditions of a typical PCR reaction (Figure 5). Following a 50-cycle reaction, the arrays synthesized on glass retained only 15 ± 3% of the initial fluorescence. After two such reactions, < 5% of the initial signal was detectable. Arrays prepared on glassy carbon and diamond exhibited greater stability. Arrays prepared on both substrates maintained ~80% of their initial fluorescence signal following initial treatment and displayed no measurable loss following the second treatment. Glassy carbon exhibited 80 ± 6% and 77 ± 5% and diamond exhibited 78 ± 5% and 74 ± 5% of their initial fluorescence levels following the first and second treatments, respectively.

**Comparison of array image quality**

Images of one-in-four arrays (a single 16 µm × 16 µm feature separated on all sides by 16 µm) were obtained using a fluorescence microscope (Figure 6). The images clearly demonstrate that there is no crosstalk between the features on any substrate. The unpolished finish of the glassy carbon surface contributes to the non-uniformity of the features at this magnification. However, this may be easily overcome by using commercially available polished surfaces. It is interesting to note, that even at this high density and small feature size, the UV dose needed to remove the NPPOC protecting group remains unchanged. The apparent increase in feature size with larger UV-dose is characteristic of the light directed in situ growth method and can be corrected for by using inverse capping (51). The polished silicon substrate behind the diamond thin film acts as a mirror, enhancing the observed fluorescence. This phenomenon is well known and is used in spotted arrays to increase signal to noise (52).

**DISCUSSION**

Because nucleic acids will not attach efficiently to untreated glass slides and because the uniformity of the
behind the thin film diamond. This may be the result of the polished silicon acting as a mirror when the diamond substrate is greater than that from either glass or glassy materials (16). The fluorescence observed from the diamond substrate is greater than that from either glass or glassy carbon. This may be the result of the polished silicon acting as a mirror behind the thin film diamond.

Figure 6. Fluorescence microscope images of a Cy3-labeled complementary DNA sequence hybridized to identical probe sequence fabricated using an increasing UV dose (measured in Joules) on glass (a), glassy carbon (b) and diamond (c). Each feature is a single pixel (16 µm × 16 µm) separated by 16 µm. The fluorescence observed from the diamond substrate is greater than that from either glass or glassy carbon. This may be the result of the polished silicon acting as a mirror behind the thin film diamond.

slide surface is critical to the quality and reproducibility of arrays, glass slides are typically modified using silane chemistries. However, the susceptibility of siloxane linkages to hydrolysis under standard conditions and the increase in the rate of hydrolysis at elevated temperatures and at basic conditions is well known (53–55). Typically, silanization is used to introduce aldehyde, amino or polylysine groups to the surface.

A similar flexibility in surface functionality can be achieved on carbon substrates by using an alkene containing the desired functional group (38,39,56,57). The carbon–carbon covalent bonds within the carbon substrate and between the substrate and the linker moiety are not susceptible to hydrolysis (54). The high stability of DNA arrays fabricated on carbon substrates, compared to their glass counterparts, reflects this fact. This increased stability of DNA arrays is important for any application where it is desirable to employ higher temperatures, extended reaction times, or basic pH conditions. Such applications include solid-phase-PCR (50,58) and surface invasive cleavage reactions (57,59–61). The stability of carbon substrates also permits their use in serial hybridization–dehybridization cycles. Arrays prepared on glass exhibit a loss of fluorescence with each subsequent chemical dehybridization cycle, while the fluorescence signal obtained from carbon substrates does not follow this trend.

The hybridization densities as measured by wash-off and collection on in situ prepared DNA arrays are lower than the densities measured when pre-synthesized oligonucleotides are immobilized on gold-thiol self-assembled monolayers (SAMs). The reduced number of hybridizable oligonucleotides per square centimeter is most likely an artifact of the in situ synthesis process. Steric hindrance due to the close proximity of the growing strand to both the surface and adjacent strands reduces the efficiencies of all steps in the synthesis process resulting in truncated DNA strands (62). While the efficiencies of coupling between NPPOC phosphoramidites and the DMT-protected phosphoramidites are comparable (14,63), post-synthesis purification for removal of truncated strands is not possible when using in situ synthesis methods. As such, the absolute number of full-length probes available for hybridization will be higher for arrays spotted on alkane-thiol SAMs. The greater the percentage of proper probe sequences, the greater the number of targets that can be subsequently captured.

This work has focused on two types of carbon substrates, glassy carbon and diamond, as materials for in situ DNA arrays. The choice of glassy carbon was based upon the commercial availability of the material from a number of vendors in either a polished or an unpolished state. Diamond deposited on silicon was included in this study to permit comparisons between this work and previously reported immobilizations of biomolecules (38,40,57). The principles established are applicable to other carbon-based materials (amorphous carbon films, nanotubes, nanorods) and carbon-rich materials (silicon carbide).

We have demonstrated that in situ photolithographic synthesis of DNA arrays on non-glass substrates is readily achievable and that these arrays offer superior stability compared to their glass counterparts under a variety of conditions. This technology enables both the repeated reuse of DNA arrays, and the use of DNA arrays for applications involving high temperatures and extremes in pH not previously accessible.

ACKNOWLEDGEMENTS

The authors thank Professor Robert Hamers for useful discussions relating to carbon surfaces, and Dr James Butler for the generous gift of the diamond substrates; Dr Kathryn Richmond for numerous discussions on DNA and synthesis quality; Omar Negrete for fluorescence microscopy assistance; Kurt Heindrich for engineering assistance and DongGee Hong for software support. This work was funded by the National Institutes of Health (R01HG02298, R01HG003275, R01EB00269), MRL is supported by the UW Technology Interface Fund. Funding to pay the Open Access publication charges for this article was provided by NIH R01HG02298.

Conflict of interest statement. None declared.

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