

Correction

ENGINEERING

Correction for “Inkjet-printed point-of-care immunoassay on a nanoscale polymer brush enables subpicomolar detection of analytes in blood,” by Daniel Y. Joh, Angus M. Hucknall, Qingshan Wei, Kelly A. Mason, Margaret L. Lund, Cassio M. Fontes, Ryan T. Hill, Rebecca Blair, Zackary Zimmers, Rohan K. Achar, Derek Tseng, Raluca Gordan, Michael Freemark, Aydogan Ozcan, and Ashutosh Chilkoti, which was first published August 7, 2017; 10.1073/pnas.1703200114 (*Proc Natl Acad Sci USA* 114:E7054–E7062).

The authors note that the following statement should be added to the Acknowledgments: “Funding support was also received from Elrha’s Humanitarian Innovation Fund (HIF)—a grant-making facility (funded by aid from the UK Government and the Swedish International Development Agency) supporting organizations and individuals to identify, nurture, and share innovative and scalable solutions to the most pressing challenges facing effective humanitarian assistance.”

www.pnas.org/cgi/doi/10.1073/pnas.1715292114

CORRECTION



Inkjet-printed point-of-care immunoassay on a nanoscale polymer brush enables subpicomolar detection of analytes in blood

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Edited by Rebecca R. Richards-Kortum, Rice University, Houston, TX, and approved July 14, 2017 (received for review February 27, 2017)

The ELISA is the mainstay for sensitive and quantitative detection of protein analytes. Despite its utility, ELISA is time-consuming, resource-intensive, and infrastructure-dependent, limiting its availability in resource-limited regions. Here, we describe a self-contained immunoassay platform (the “D4 assay”) that converts the sandwich immunoassay into a point-of-care test (POCT). The D4 assay is fabricated by inkjet printing assay reagents as microarrays on nanoscale polymer brushes on glass chips, so that all reagents are “on-chip,” and these chips show durable storage stability without cold storage. The D4 assay can interrogate multiple analytes from a drop of blood, is compatible with a smartphone detector, and displays analytical figures of merit that are comparable to standard laboratory-based ELISA in whole blood. These attributes of the D4 POCT have the potential to democratize access to high-performance immunoassays in resource-limited settings without sacrificing their performance.

nanoscale | nonfouling | polymer brush | inkjet printing | point of care

Diagnostic assessment of protein biomarkers plays an essential role in modern medical practice, and its availability has a considerable impact on clinical evaluation and decision making in human health and disease (1). Diagnostic biomarkers are not only used for identifying disease in individual patients but also for developing treatment strategies, tracking treatment response, monitoring recurrence, conducting clinical trials, and performing epidemiological analysis (2). However, shortages in sophisticated laboratory resources, highly trained personnel, and modern facilities (with clean water, cold storage, and reliable power) have hampered the discovery and use of diagnostic and therapeutic biomarkers in the developing world (3). The mainstay approach for sensitive and quantitative detection of protein biomarkers is the ELISA. In developed countries, highly sensitive ELISA and ELISA-like (4) sandwich immunoassay technologies are readily accessible in centralized facilities and are performed by trained operators or even automated to handle the multistep workflow. Much of the workflow is directed toward reducing biomolecular noise due to nonspecific binding of proteins and other components in complex biological fluids that lower signal-to-noise ratio (SNR). Depending on the assay, steps can include sample pre-processing, liquid transfer, blocking, incubation, and wash steps, in addition to data acquisition and analysis with bulky instrumentation. Taken together, these characteristics represent major barriers to obtaining highly sensitive quantitation of protein biomarkers in limited-resource settings (LRSs) (3). Furthermore, these barriers can also delay treatment, as test results from laboratory-based immunoassays may not always reach healthcare providers and patients in LRSs quickly enough to impact critical clinical decisions (1). To address these concerns, we sought to design and implement a broadly applicable protein

biomarker detection platform with performance comparable to resource- and personnel-intensive technologies such as ELISA while retaining the many attractive features (i.e., low cost, portability, and ease of use) of “passively” driven platforms such as lateral flow immunoassays (LFIAs) (5), paper-based diagnostics (PBDs) (6), and passive microfluidics (PMFs) (7).

In our previous work, we fabricated Ab-based microarrays on nanoscale poly(oligo(ethylene glycol) methacrylate) (POEGMA) polymer brushes that were capable of femtomolar detection of analytes directly from complex biological milieu (8). The Ab arrays were directly spotted onto POEGMA brushes that were grown from glass by surface-initiated atom transfer radical polymerization (SI-ATRP) (8, 9). The POEGMA’s ability to resist nonspecific adsorption of proteins greatly improved SNR by reducing “biomolecular noise,” translating to femtomolar limit of detection (LOD) of protein analytes even from whole blood and serum (8, 10, 11). The observation that Ab microarrays could be directly spotted and noncovalently immobilized onto POEGMA coatings, and that spots of Abs dried and captured within the POEGMA brush retained their activity even after drying and ambient storage, greatly simplified assay fabrication

Significance

Sensitive quantitation of protein biomarkers plays an important role in modern clinical decision making. This work introduces an inkjet-printed assay platform built on a nonfouling, nanoscale polymer brush, which eliminates nonspecific binding, the largest source of noise in surface-based assays. The assay goes to completion after adding a drop of blood (with no additional reagents or mixing), and the assay can be read with a smartphone-based detector. This technology is significant because it enables high-performance diagnostic testing in blood with minimal infrastructural requirements. Furthermore, its fully printed nature makes it highly customizable and thus broadly applicable to a wide range of diagnostic targets.

Author contributions: D.Y.J., A.M.H., Q.W., K.A.M., C.M.F., R.T.H., R.B., Z.Z., R.K.A., D.T., R.G., M.F., A.O., and A.C. designed research; D.Y.J., A.M.H., Q.W., K.A.M., M.L.L., R.B., Z.Z., R.K.A., D.T., R.G., M.F., and A.O. performed research; A.O. contributed new reagents/analytic tools; D.Y.J., A.M.H., Q.W., K.A.M., M.L.L., C.M.F., R.T.H., R.B., Z.Z., R.K.A., D.T., R.G., M.F., A.O., and A.C. analyzed data; and D.Y.J., A.M.H., Q.W., M.L.L., R.G., M.F., A.O., and A.C. wrote the paper.

Conflict of interest statement: The underlying technology of the D4 was developed by A.M.H. and A.C. and acquired by Immucor Inc. in 2014.

This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703200114/-DCSupplemental.

and subsequent storage of the microarrays. Significantly, the POEGMA-based microarrays also decreased the time needed to perform a sandwich immunoassay by eliminating the need for blocking steps and reducing the number of rinsing steps. Like most other immunoassays, however, this approach still required separate incubation steps for labeling with detection reagents to achieve a quantifiable signal. We therefore sought to build upon this body of work and the robust design concepts of existing “point-of-care test” (POCT) technologies to develop a quantitative and sensitive platform that stores all necessary reagents so that the assay is ready for readout following direct addition of blood or serum, while minimizing user intervention.

Here, we report a quantitative, self-contained, multiplexable immunoassay (the “D4 POCT”) that stores all necessary capture and detection reagents “on-chip” in a dry state, shows durable stability without cold storage, and detects analytes with sensitivity comparable to ELISA directly from whole blood without sample preprocessing.

Results

Design of the D4 Immunoassay. The challenge we faced in redesigning an immunoassay on POEGMA brushes as a POCT, and one that is common to most POCTs, is how to spatiotemporally separate the capture and detection reagents, and how to time the precise sequence of events upon introduction of the analyte into the assay to yield a readout. Whereas recent work in PBDs that use highly innovative and robust sequential delivery techniques has been shown to be effective (12–15), our approach was to instead design a passive POCT that uses lateral diffusion in the polymer brush as the temporally programmable mixing principle. The architecture of the D4 assay consists of a glass chip with a ~50-nm-thick POEGMA coating that contains two types of microspots that are inkjet-printed on the polymer brush: “stable” spots of capture antibodies (cAb) and “soluble” spots of detection reagents (Fig. 1). Here, the capture and detection reagents are placed in very close proximity to one another (several hundred micrometers), unlike LFIA in which reagents must travel larger distances. The detection reagent spots are soluble, as they consist of a mixture of fluorescently labeled detection Abs (dAbs) and excipient (such as PEG or trehalose) to enable dissolution upon liquid exposure (Fig. 1 *A* and *B*). The D4 assay is so named because of the chain of events that drive the assay to completion upon addition of a drop of blood (Fig. 1 *A–D*): (*A*) dispense blood onto chip, (*B*) dissolution of “soluble” detection reagent spots, (*C*) diffusion of analyte-bound dAb across surface and binding to respective cAb spots, and (*D*) detection of binding event by fluorescence imaging. An overall schematic of the fabrication of the D4 assay chip is shown in Fig. 1*E* and consists of the following sequential steps: (*i*) SI-ATRP to grow ~50-nm-thick POEGMA brushes on the glass surface, (*ii*) followed by noncontact inkjet printing of the cAb from buffer and a fluorescently labeled dAb with a molar excess of an excipient, typically PEG.

D4 Detection of Human IgM/IgG from Fingerstick. We first sought to test the concept of the D4 POCT and chose the detection of human IgG and IgM as a test case, as both analytes are present at high concentration in blood and should thus be detected by the assay. Furthermore, detection of IgG and IgM in the same assay also allowed us to assess the feasibility of carrying out a multiplexed D4 assay. To test this concept, a POEGMA-coated glass slide was stamped with a wax grid to confine the sample to the active area of the chip containing an inkjet-printed Ab array (Fig. 2*A*). The detection mixture was printed as three outer rings of soluble dAb spots containing a mixture of mouse Cy-5-anti-human-IgG and/or mouse Cy-5-anti-human-IgM (dAbs with a different epitope against human IgG and IgM than the cAb), with linear PEG (molecular weight 116,000), an excipient that

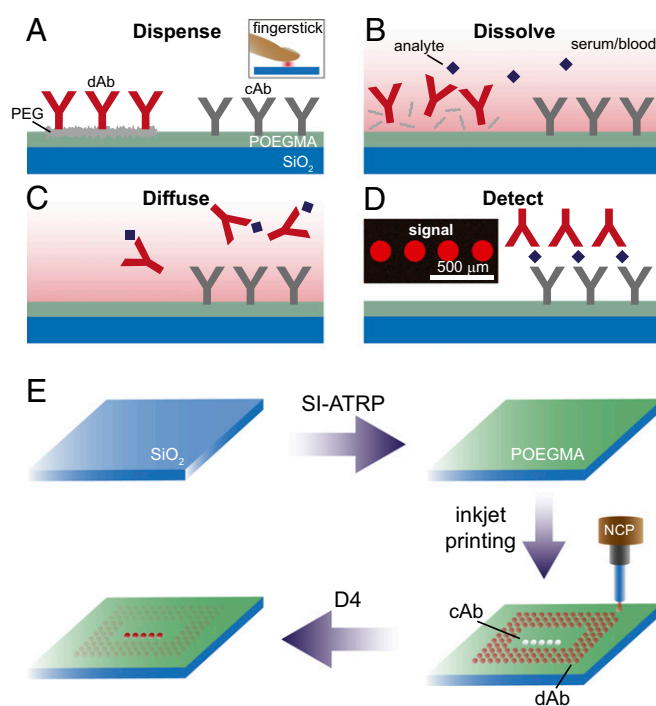


Fig. 1. D4 immunoassay on POEGMA brushes. (*A*) “Stable” spots of cAb and “soluble” spots of the fluorescently labeled dAb are printed onto the POEGMA brush. Whole blood or serum is dispensed directly onto the chip. (*B–D*) The sequence of events after addition of blood or serum is as follows. (*B*) “Soluble” dAb dissolves and binds to analyte. (*C*) These complexes diffuse and bind to their respective cAb spots, and subsequently (*D*) generate a quantifiable fluorescent signal. (*E*) D4 chip fabrication. Glass chips are coated with POEGMA with SI-ATRP. The cAb and detection reagents are spotted onto the surface with noncontact inkjet printing (NCP). After drying, chips are ready for use.

was added to enable dissolution of the dAbs upon contact with blood, and heparin to prevent coagulation of blood. The inner 4×4 array contains spots of “stable” cAb. Rows 1 and 4 are an anti-mouse cAb (positive control), row 2 is an anti-human IgG cAb, and row 3 is an anti-human IgM cAb.

Representative steps performed by a user are shown in Fig. 2 *B–D*. First, a drop of blood from a finger stick (Fig. 2*B*) is applied directly to the microarray and contained within the hydrophobic corral (Fig. 2*C*). After a predetermined incubation period (5 min for this experiment), the surface is rinsed with ~1 mL wash buffer from a squeeze bottle, which displaces the loosely bound blood cells and proteins (Fig. 2*D*). Interestingly, the blood flows to the margins and binds to the hydrophobic corrals, as seen from the red color around the margins in Fig. 2*D*, but is completely removed from the non-fouling POEGMA surface. The slide is then imaged with a fluorescence detector. The output of the D4 from a fluorescent scanner is shown in Fig. 2 *E–J*. Before exposure to blood, the inner 4×4 cAb array has no intrinsic fluorescence (Fig. 2*E*). Negative control experiments in which the D4 chip was incubated with either PBS (Fig. 2*F*) or whole chicken blood (Fig. 2*G*) show that only the positive-control spots (rows 1 and 4) generate signal, while middle rows 2 and 3 (specific for human IgG and IgM, respectively) show no fluorescence. Upon incubation with human blood with an array that is printed with a detection mixture of both Cy5-anti-human IgM and Cy5-anti-human IgG dAbs (Fig. 2*H*), rows 2 and 3 show positive fluorescence signal by “sandwiching” circulating human IgG (row 2) and IgM (row 3) analytes between a cAb and the Cy5-labeled dAb specific to each analyte. In contrast, when the outer dAb spots only contain Cy5-anti-human IgM dAb (Fig. 2*I*), only row 3 (IgM cAb) is

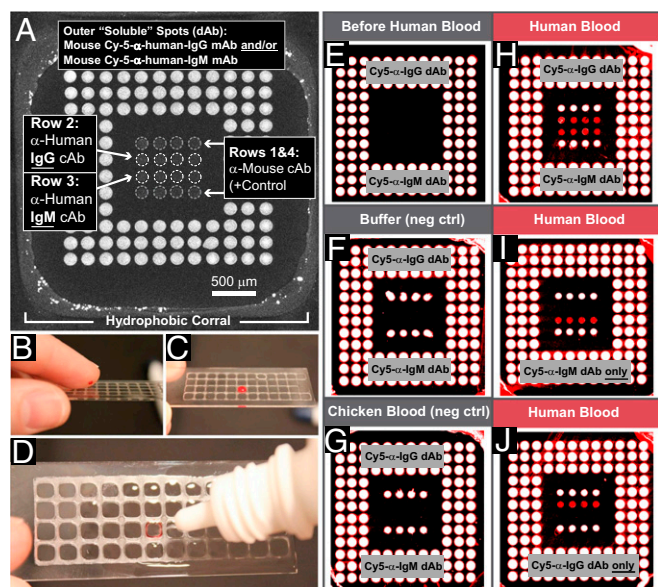


Fig. 2. D4 detection of human IgG and IgM from fingerstick blood. (A) D4 microarray within a single wax corral. Outer “soluble” dAb spots: murine Cy-5-anti(α)-human-IgG dAb and/or murine Cy-5- α -human-IgM dAb. Inner 4×4 “stable” cAb spots: α -mouse cAb as positive control (rows 1 and 4), α -human IgG cAb (row 2), or α -human IgM (row 3). (B–D) Steps performed by use. (B) Application of fingerstick blood, (C) incubation for 5 min, and (D) washing with 1 mL of buffer from squirt bottle. (E–I) Results of D4 assay. (E) Before liquid exposure. (F and G) After exposure to PBS and whole chicken blood. (H–J) Results after adult human fingerstick blood (5 min). Each outer spot contains a mixture of PEG excipient, heparin, and (H) both Cy5- α -human-IgG and Cy5- α -human-IgM dAbs, (I) only Cy5- α -human-IgM dAb, or (J) only Cy5- α -human-IgG dAb.

visible but not row 2 (IgG cAb). The converse is true when the outer spots only contain Cy5-anti-human IgG dAb (Fig. 2I).

In summary, this proof-of-concept experiment demonstrated that printing both cAb and dAb together on PEOGMA coatings greatly simplifies the sandwich immunoassay by eliminating the multistep procedures typically used in standard ELISA. It also suggested that multiplexed detection should be possible with the D4 assay, a desirable feature that has been historically difficult with LFIA (16).

Quantitative Single-Analyte D4. The concentration of total Ig is relatively high in whole blood [approximately in the micromolar range (17)], so we next sought to examine the quantitative response of the D4 assay for analytes that occur at lower concentrations in complex biological matrices such as serum and whole blood. Fig. 3A shows representative image data of a D4 assay for the cytokine IL-6 exposed to whole chicken blood with and without spiked analyte (right and left panels, respectively). Similar to the array design in Fig. 2, soluble spots of anti-IL-6 dAb are printed as outer rings surrounding centrally located capture spots of anti-IL-6 cAb. These anti-IL-6 cAb spots were printed alongside spots of vehicle control (PBS) and positive-control capture spots (labeled “ctrl”) comprised of anti-dAb Abs targeting the Fc portion of dAbs. These positive-control spots not only indicate whether dAbs were successfully localized to the active area of the assay but also help quantitatively correct for interassay variation as calibration spots (SI Appendix, Fig. S1). The images in Fig. 3A show minimal background signal, due to the nonfouling polymer brush, near-zero signal from anti-IL-6 cAb spots in the absence of analyte, and brightly fluorescent anti-IL-6 cAb spots in the presence of human IL-6-spiked blood. As expected, D4 signal was absent from vehicle control and brightly

fluorescent in positive-control spots in both cases. By measuring the fluorescence emission from cAb spots across a range of different concentrations in IL-6-spiked serum and whole blood (Fig. 3B), we obtained the dose–response curves shown in Fig. 3C and D. For this IL-6–D4 assay, we calculated that the LODs in serum and blood were approximately 6.3 pg/mL and 10.9 pg/mL, which correspond to ~ 310 and 536 fM, respectively, with a dynamic range spanning greater than three orders of magnitude [details for calculating figures of merit (FOM) are provided in Materials and Methods].

The D4 is a flexible platform for rapid immunoassay development. By inkjet printing spots of stable cAb and soluble dAb for various targets onto PEOGMA-coated chips in a layout like that of Fig. 3A we were able to develop D4 chips for a variety of different markers. To demonstrate this versatility, in SI Appendix, Fig. S2 we show the performance of D4 immunoassays pertinent to oncology [prostate-specific antigen (PSA) and alpha-fetoprotein (AFP)], endocrinology (leptin), cytokine profiling (TNF α and IL-6), cardiology [B-type natriuretic peptide (BNP)], and infectious disease (HIV p24), in both serum and whole blood. The quantitative FOM for each analyte are summarized in Table 1. The data show that for each analyte tested the D4 is quantitative and sensitive with LODs that range from approximately 6.3 to 113 pg/mL, dynamic ranges between 2.3 to 3.9 orders of magnitude, and acceptable intra- and interassay coefficients of variation (COV_{intra} and COV_{inter} ; no greater than 15% and 20%, respectively). While the assays in Table 1 used incubation times of 90 min to maximize sensitivity, we observed that much shorter times can be used for sensitive assay readout. To examine the effect of incubation time on LOD, we carried out a leptin–D4 assay for incubation times ranging from 15 min to 90 min and found that the LOD only increased to 57 pg/mL for a D4 assay carried out for 15 min compared with 38 pg/mL for an assay carried out for 90 min (SI Appendix, Fig. S3).

An important design feature that we also considered in our implementation of the D4 was accounting for variability in factors such as sample volume and dissolution efficiency of soluble dAb, which may impact signal quantitation. Our approach to do so is described in greater detail in SI Appendix. Briefly, however,

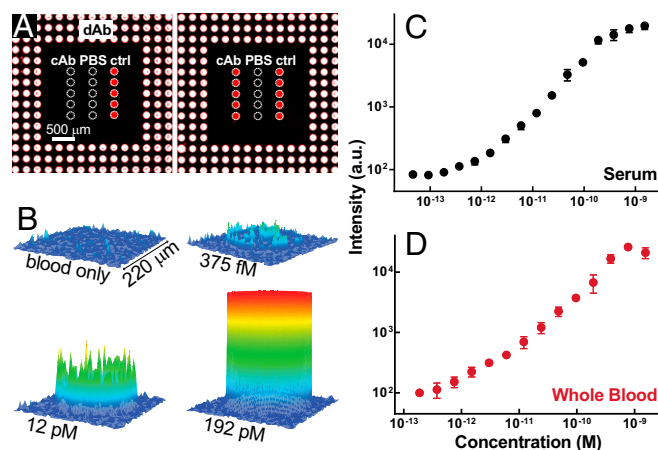


Fig. 3. Representative D4 against IL-6 in whole serum and blood. (A) D4 image data. Spots of Cy5-dAb against human IL-6 are printed around spots of anti-IL-6 cAb. Spots of PBS and anti-dAb Abs printed nearby serve as negative and positive controls, respectively. All spots are artificially outlined by a dashed white line to aid visualization of spot locations. Images show arrays exposed to whole chicken blood alone (Left) and blood spiked with IL-6 (Right). (B) Spatial intensity plots of fluorescence from individual cAb spots at various concentrations of IL-6 spiked in whole chicken blood. (C and D) Dose–response curves of IL-6 in (C) calf serum and (D) chicken blood. Each data point represents mean \pm SD from three separately run D4 arrays. LODs in serum and blood were 6.3 and 10.9 pg/mL, respectively.

Table 1. D4 assay FOM

Analyte	LOB, pg/mL	LOD, pg/mL	COV _{intra} %	COV _{inter} %	DR, log ₁₀
AFP					
Calf serum	20.1	47.5	4.7	14.5	2.7
Chicken blood	30.0	58.9	6.3	13.5	2.8
PSA					
Calf serum	54.4	82.8	5.5	10.5	2.7
Chicken blood	51.0	112.9	8.0	11.0	2.5
Leptin					
Calf serum	10.3	38.2	6.4	8.2	3.2
Chicken blood	10.4	44.3	4.8	13.4	3.1
BNP					
Calf serum	2.1	25.2	10.8	19.1	3.7
Chicken blood	8.7	28.2	8.9	15.2	3.3
IL-6					
Calf serum	3.8	6.3	7.6	10.0	3.4
Chicken blood	3.7	10.9	7.1	14.1	3.1
TNF α					
Calf serum	2.0	11.1	9.8	19.9	3.0
Chicken blood	24.3	54.2	7.4	18	2.3
HIV p24					
Calf serum	4.8	11.0	9.9	13.3	3.9
Chicken blood	3.1	16.5	9.1	15.6	3.4

the effect of these variations can be eliminated by incorporating positive control calibration spots comprised of anti-dAb Abs that target the Fc portion of dAbs that are printed in the center of the D4 arrays alongside anti-analyte cAb spots, as shown in the image data of Fig. 3A and *SI Appendix, Fig. S1A*. The data in *SI Appendix, Fig. S1* show a significant reduction in interassay variability upon normalization of the cAb spot intensity to the fluorescence intensity of control spots, and the normalized signal intensities are consistent across a range of sample volumes from 50 to 150 μ L.

Quantitative Multianalyte D4. We next assessed the performance of the D4 assay to simultaneously quantify multiple biomarkers using an approach similar to that shown in Fig. 2. Printing spatially distinct capture spots on a 2D surface enables the detection of multiple targets from a single chip with the same fluorescent reporter. As a proof-of-concept demonstration, we developed duplexed assays against cytokine markers TNF α and IL-6 (Fig. 4) and cancer markers AFP and PSA (*SI Appendix, Fig. S4*). Fig. 4A shows an image of the multiplexed D4 against cytokines TNF α and IL-6 after incubation with analyte-negative chicken blood and indicates the location of capture spots and Cy5-labeled detection reagents. In a format similar to that of the single-analyte assays shown earlier, labeled detection reagents containing dAb for TNF α and IL-6 were printed in the region surrounding cAb spots.

As shown in Fig. 4B, performing the D4 in whole blood spiked with a mixture of both TNF α and IL-6 shows detectable fluorescence at both cAb spots for each analyte, in a dose-dependent manner, similar to that of the single-analyte assays shown in *SI Appendix, Fig. S2*. In contrast, Fig. 4C and D show that when the multiplexed assays are exposed to either TNF α or IL-6 alone only cAb spots specific to each respective analyte show dose-dependent fluorescence. Similar multiplexed data for AFP and PSA are shown in *SI Appendix, Fig. S4*.

Pilot Clinical Study with D4. We next sought to test the D4 in human patients using a biomarker that would be useful for POC applications. From the panel of analytes shown in Table 1 we elected to test the performance of a D4 against leptin in a clinical setting. This choice was motivated by a recent study in Uganda by investigators at Duke University and Mulago Hospital which determined that a low serum leptin level (even below 50 pg/mL)

is a major biochemical risk factor predicting infant mortality due to malnutrition (18). The authors of this study suggested that leptin measurements could be used to identify and provide targeted treatment to malnourished children at highest risk of death. Unfortunately, many low-resource settings lack the infrastructure to perform hormone assays at such sensitivities and require blood specimens to be transported to larger facilities, which delays potentially life-saving treatment. We therefore speculated that the D4 may provide a useful diagnostic alternative to hormone assays carried out in a centralized laboratory by allowing sensitive and quantitative on-site leptin measurements.

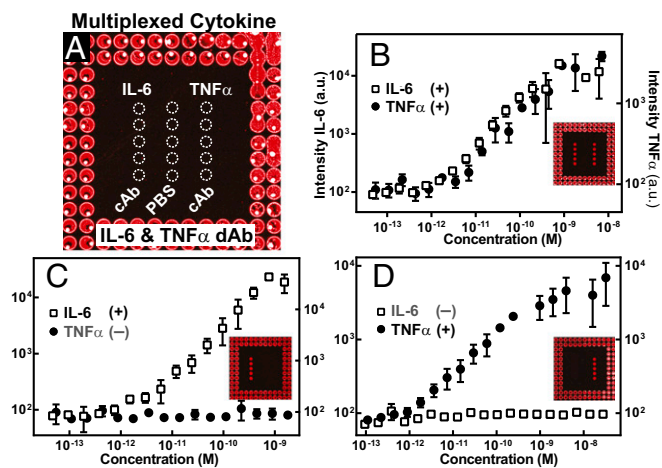


Fig. 4. Multiplexed assays against cytokine markers in whole blood. Fluorescent detection reagents against both analytes are coprinted as outer spots. Spots of cAb against IL-6 and TNF α are printed in the center of the array. (A) D4 image after incubation with whole chicken blood alone (without analyte). For clarity, spots for cAb and PBS are indicated with white dotted lines. (B) Dose-response curves after exposure to whole chicken blood spiked with a mixture of both IL-6 and TNF α analytes at varying concentrations. (C and D) Dose-response curves when assay exposed to varying concentrations of (C) IL-6 only or (D) TNF α only. Insets for panels B–D show D4 image data. Each data point represents mean \pm SD from three separately run D4 arrays.

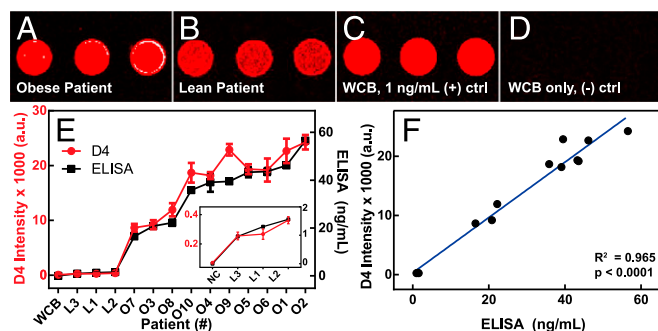


Fig. 5. Pilot clinical studies using D4 to measure leptin levels in pediatric patients. (A–D) Representative D4 imaging of whole blood specimens from an (A) obese patient and (B) lean patient immediately after blood draw. For comparison, D4 data of (C) whole chicken blood (WCB) spiked with 1 ng/mL human leptin (D) and human leptin-negative chicken blood alone are also shown. (E) D4 fluorescence readout compared with results from standard sandwich immunoassay in all patients. D4 data are shown in red and sandwich immunoassay data in black. (Inset) Zoomed graph of lean patients alone (NC = negative control serum). (F) Plot showing correlation between D4 versus sandwich immunoassay ($R^2 = 0.965$; $P < 0.0001$, two-tailed t test).

As proof of concept, we conducted a small IRB-approved pilot study at Duke University Medical Center (DUMC) comparing serum leptin levels detected by the D4 assay versus clinical ELISA from 3 lean [body mass index (BMI) $<10\%$] and 10 obese (BMI 83.9–99.58%) pediatric patients. We selected this patient population to clinically test the D4 assay across a wide range of leptin levels, as serum leptin levels are proportional to adipose mass (19, 20). Our pilot studies demonstrated high concordance between D4 and clinical ELISA (Fig. 5). Fig. 5A shows D4 images from a representative obese and lean patient in whole blood, immediately after blood draw. The obese patients exhibited higher fluorescence intensities (indicating higher leptin levels) than lean patients (Fig. 5B), which is consistent with known adipocytokine physiology. Significantly, we found that the leptin levels measured by D4 correlated strongly with those measured by clinical ELISA performed in parallel in a central laboratory at DUMC across all 13 patients ($P < 0.0001$) (Fig. 5E and F).

The readout of D4 microarray chips shown thus far was obtained using a sensitive table-top fluorescence scanner to assess the sensitivity of the D4 assay. While this approach would allow for POC testing in a peripheral laboratory near or attached to a clinic in LRSs (21), we recognize that a table-top scanner is too burdensome for use in the field. To address this issue, we next investigated the feasibility of portable fluorescence imaging of the D4 assay using a mobile phone-based fluorescence microscope (Fig. 6 and *SI Appendix*, Fig. S5). This apparatus is a compact and cost-effective imaging system that uses an external lens, in addition to the existing lens of the mobile phone camera (22, 23). The illumination is introduced at an oblique illumination angle of $\sim 75^\circ$ to increase the SNR of the acquired fluorescence images on the phone.

These experiments imaged leptin-D4 assays in both simulated and patient samples. For comparison, we also imaged the same set of D4 arrays (to maintain consistency) with the table-top scanner in the usual fashion. The fluorescence images in Fig. 6A and B depict representative cAb spots of leptin-D4s (with and without analyte) using the table-top scanner and the mobile phone-based imager, respectively. In both cases, the fluorescence readout behaves as expected for leptin-spiked and leptin-deficient serum (left and right image panels), with good SNR. The dose–response curves using a dilution series of leptin-spiked calf serum are shown in Fig. 6A and B. Both scanner and mobile phone-based imaging modalities showed quantitative, dose-dependent fluorescence intensities. In this experiment,

the scanner was more sensitive than the mobile phone, with LODs determined to be 0.037 ng/mL and 0.71 ng/mL, respectively. We next imaged a set of leptin-D4 arrays against clinical specimens obtained from obese patients in our pilot clinical study. The D4 readouts from the scanner (red trace) and from the mobile phone platform (blue trace) are shown in Fig. 6C. In both cases, as expected the data show good correspondence with ELISA values (dashed black trace).

Transitioning to the mobile phone-based detection platform reduced detection sensitivity and interassay consistency, as seen by the higher LOD and larger error bars with the mobile phone device. Nevertheless, these proof-of-concept studies demonstrate the feasibility of merging D4 assay technology with compact, field-portable, cost-effective, and easy-to-use mobile phone-based detection platforms. As mobile phone detector technology, computational imaging, and sensing approaches continue to evolve we expect that the fluorescence collection efficiency and hence sensitivity of these portable, low-cost detectors will rival table-top fluorescence scanners. Despite the lower sensitivity of this first-generation mobile phone detector, the field portability of mobile phone-based imaging provide a detection strategy that is well-matched to complement the robustness of the D4 assay technology with FOM that are sufficient for many clinical applications.

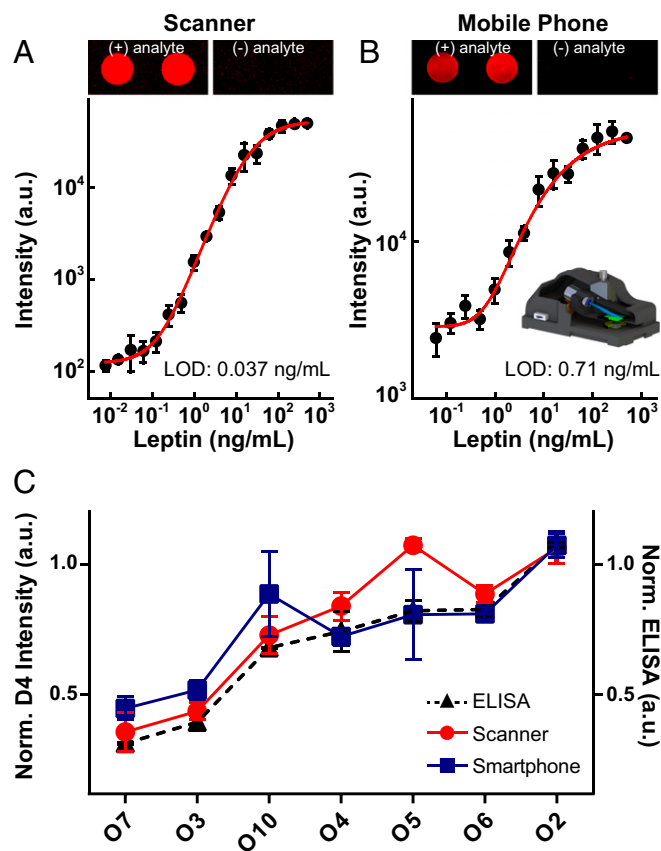


Fig. 6. Mobile phone-based imaging of D4 arrays. (A and B) Representative image data of D4 microspots and dose–response curves in calf serum spiked with leptin analyte acquired by a (A) benchtop scanner and (B) our mobile phone-based fluorescence microscope. Five-parameter logistic fit shown as red curve. (B, Inset) Illustration of the mobile phone attachment for fluorescence imaging and quantification of D4 arrays. (C) Testing of obese patient serum with D4 using a scanner (red trace) versus the mobile phone microscope (blue trace), and comparison with ELISA results (dashed black trace). Normalized D4 data (scanner, phone) plotted on the left axis, and normalized ELISA data plotted on the right axis.

Storage Stability of D4. Because use of the leptin D4 POCT in LRSs would necessitate long-distance shipment of assay materials to sites around the world, an important issue that needed to be addressed is the need for a “cold chain” and the storage stability of D4 chips. We hence measured the performance of D4 chips targeting leptin at different time points and under elevated temperatures after chip fabrication. Fig. 7 compares dose–response curves of leptin-D4s exposed to analyte-spiked calf serum under different storage conditions. In Fig. 7A, vacuum-sealed packets containing leptin-D4 chips were stored 1–92 d under ambient conditions. The data showed no significant difference in assay performance. Likewise, we also observed minimal difference in assay performance when vacuum-sealed chips were stored at 45 °C for up to 120 h (5 d) (Fig. 7B).

Discussion

The D4 POCT offers a promising platform for democratizing access to sensitive and quantitative assessment of diagnostic protein biomarkers. We first showed proof-of-concept application of multiplexed detection of human immunoglobulins. We observed ELISA-like performance across a wide range of protein analytes when assayed in unmodified serum and blood and showed the feasibility of quantification for multiple analytes from the same chip. Our pilot validation studies showed high concordance between D4 and ELISA in human patients, suggesting that this platform may ultimately be translatable to the clinical setting. Next, we introduced the feasibility of combining D4 chip technology with sensitive and cost-effective mobile phone readout for truly distributed use of the platform. Finally, the D4 chips showed durable storage stability, as evidenced by acceptable assay performance following exposure to heat or prolonged storage.

Much work has been previously done toward developing POCT platforms for LRSs. An “active” approach is to incorporate a fluid handling system where the reagents are stored on the device separately and then mixed and washed using pumps and valves, or alternatively by mechanisms that are conceptually similar but with smaller microfluidic devices (7, 24–27). This pioneering design strategy has seen commercial success in some embodiments (e.g., Abbott i-Stat) and shown encouraging field performance in others (26, 27). However, such systems have many parts, which increases the complexity and cost, and the finite possibility of malfunction from each individual component can limit the robustness of the device.

To date, “passively” driven immunoassays have experienced the most success as POCTs for protein analyte detection. LFIAs—the most widely used embodiment in LRSs—use nitrocellulose strips impregnated with dried capture/detection reagents that are passively collocated by applying liquid; this design permits a low-cost, rapid approach to protein detection in a user-friendly manner (fluid application with or without premixing, sometimes followed by buffer addition step and/or signal amplification) (5, 28). Despite

its strengths, the LFIA is not a substitute for the ELISA for two major reasons. First, LFIAs provide semiquantitative or qualitative readouts (in the absence of a dedicated reader), while conventional ELISA provides quantitative results with high accuracy and precision. Second, most LFIAs are not as sensitive as ELISA (29–31), as more antigen-antibody (Ag–Ab) interactions are necessary to generate a positive signal. This leads to certain clinical scenarios in which LFIA technology is unable to meet the required sensitivity to be practically useful (32). Furthermore, PBDs have been groundbreaking in further evolving passively driven devices for use in LRSs and offer many advantages over existing POCTs for their low cost, disposability, and ability to be assembled into multidimensional structures (33, 34). While a frequently encountered shortcoming with PBDs has been limited sensitivity, several innovative signal amplification techniques that reduce detection limit without significantly compromising usability have been described, most notably those using paper networks for automated sequential reagent delivery (12, 15) and enzymatic amplification (35). Such advances have made PBDs a promising platform for “lab-on-a-chip” diagnostics; however, more work remains toward addressing issues such as controlling the consistency of paper matrices, the underlying need for the sequential delivery of reagents and/or washing, challenges with multiplex analysis, and further improving detection sensitivities (34, 36). In contrast, capillary-driven PMFs (e.g., MBio and Philips Minicare) have overcome challenges in sensitivity faced by LFIAs and PBDs while retaining ease of use, with some platforms reaching subpicomolar LODs within 15 min (7, 37–42). By eliminating “active” elements (e.g., pumps), PMFs reduce the complexity and cost of microfluidic-based designs and significantly reduce instrumentation footprint compared with standard assays. While PMFs show great promise, challenges remain in further reducing instrumentation footprint and/or lowering cost of readers to fulfill the accessibility and sustainability requirements that are critical for successful dissemination and implementation in LRSs.

Our contribution toward democratizing accesses to clinical diagnostics—the D4—builds upon, but also departs significantly from, the accomplishments of existing “passive” POCT designs such as LFIAs, PBDs, and PMFs. The POEGMA interface is the critical element responsible for the high performance and simplicity of the D4. In the D4, the POEGMA brush is nonfouling in its hydrated state. The hydrated POEGMA brush virtually eliminates nonspecific binding of cells, proteins, and other biomolecular noise (8, 9, 43), yielding high SNRs even at low analyte concentration in complex matrices such as blood and obviates the need for microfluidic separation (preprocessing) of cells.

In the dry state, however, the POEGMA brush readily entraps Abs that are inkjet-printed into the polymer brush. Printing cAbs onto the dry brush without excipient leads to their stable immobilization within the brush and prevents bleeding or dissolution of cAb spots upon hydration with biological matrix (8). This leads to uniform capture spot morphology, which facilitates quantification and analysis. In contrast, printing detection reagents with an excess of soluble excipient onto the dry brush creates a reservoir for assay reagents. Contact with blood or other aqueous fluid causes dissolution of the excipient in printed spots of the dAb and releases the dAb from the chip, and the released dAb freely diffuses across the surface and drives the assay to completion. The direct inkjet printing of reagents onto POEGMA brush is another key element of the D4 assay, as it allows precise control over assay geometry, spot concentration, and composition, without the need for covalent coupling steps, which simplifies assay fabrication. Like many existing “passive” designs, the D4 stores all necessary reagents on the same device, but it does so by inkjet printing capture and detection reagents in very close proximity onto a nonfouling polymer brush. This design eliminates the need to control fluid movement with fluidics or include sequential liquid transfer steps. Combined, these features

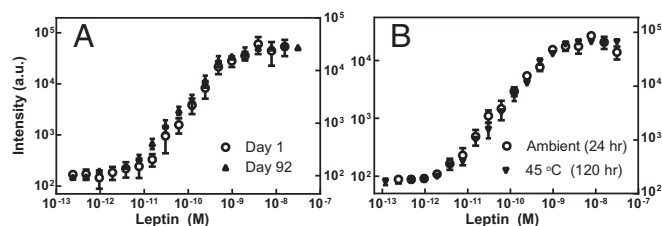


Fig. 7. Storage stability of the D4. Leptin-D4 chips were stored in vacuum-sealed packets under the conditions described and then run in analyte-spiked calf serum. (A) Dose–response curves for leptin-D4 after 1 d (○) and 92 d (■) of storage in ambient temperature show similar performance. (B) Dose–response curves for leptin-D4 after 24 h of ambient storage versus 120 h (5 d) of storage at 45 °C show similar performance.

of the D4 platform translate to a robust, self-contained, and quantitative POCT without compromising assay performance or cost.

The low-cost fabrication of the D4 and smartphone-based reader, coupled to its sensitivity and quantitative performance, may offer a scalable and sustainable route to manufacturing multiplexed immunoassays for LRSs that are both affordable and analytically robust. When produced in bulk, we estimate that D4 chips would cost <\$1 each (*SI Appendix, Table S1*) and the 3D-printed mobile phone attachment would cost <\$30 each. In terms of workflow, we envision that the D4 platform can provide a streamlined protocol for protein analyte detection in both healthcare and research settings and can expand the capabilities of both. The ability to achieve multiplexed results with ELISA-like sensitivity after a single incubation/rinse step and without blocking greatly simplifies protein quantification, reduces time to results, and ultimately lowers costs. Admittedly, creating a multiplexed test in LFIA format is also relatively straightforward as it involves spotting multiple capture reagents onto a strip or adding additional test “lines.” However, the LFIA is highly dynamic, as its constituents are constantly changing at a given location across a test strip and never fully equilibrate. Subsequently, assay readout relies on binding kinetics specifically during which reactants are sufficiently close to interact on a molecular level, and this varies for each individual test. Unless each test is isolated from the others, the same assay condition (e.g., running buffer, porous membrane, and flow rate) is applied to all tests, and one set of conditions is often not optimal for the multiple analytes of interest. While isolation of individual tests is not overly difficult and various strategies to this end are available (44), this adds to complexity of sample handling and cost. In contrast, the design of the D4 assay—by effectively colocalizing the detection and capture reagents and relying on 2D diffusion and equilibration “above” a nonfouling polymer brush—allows multiplexing without significantly increasing the assay complexity.

Difficulty in storage is a major factor restricting the implementation of laboratory-based tests in LRSs. We speculate that the storage stability of D4 is related to the high oligo(ethylene glycol) density of the POEGMA “bottle brush,” which provide a protective environment that stabilizes the printed proteins against denaturation (8, 45). To put our results into context, LFIAs are already well-suited for LRSs since they are generally stable for 1 to 2 y at ambient temperature and 6 mo at elevated temperature (45 °C) (16), and similar reagent stability has also been shown in other POCTs (46, 47). The timescales for storage stability used in our studies with the D4 (Fig. 7) were shorter than those determined for LFIAs. Thus, while encouraging, we recognize that further studies are required to characterize the storage limitations of the D4 and, if needed, modifying packing and/or printing solutions with existing methods for reagent stabilization (35, 48) to meet the storage demands of LRSs.

For patient care, we envision that this technology will provide a robust POCT that is widely applicable for the diagnosis of a disease by quantification of the levels of one or more markers for which orthogonal Ab pairs are available. In the developed world, this will bring highly sensitive immunoassays to first responders and to the bedside, while in LRSs it will allow them to simply become more routinely available. The low cost of the chips and detectors also suggests that wide dissemination of this technology is feasible, so that it would also provide new opportunities for spatiotemporal mapping and remote analysis of measurement results, which would be highly valuable for epidemiology in general. However, it is worthwhile to note that the actual need for ELISA-like sensitivities depends on the disease and biomarker, which must be considered when deciding between the D4 POCT and other robust alternatives such as LFIAs or PBDs when such sensitivities are not required.

There are several challenges that still need to be resolved. The first is the requirement to remove the blood from the chip before imaging. Because proteins do not adsorb to the POEGMA brush

surface and cells similarly do not adhere and are only settled on to the surface by gravity, removal of the blood from the chip is currently easy to perform using a buffer rinse. We acknowledge, however, that the open wash requirement is not ideal for bio-safety reasons and possible specimen contamination. The ability to read out the assay in a sealed system with an automated wash or even in the presence of blood would make this a completely hands-off POCT, which is desirable. We are currently pursuing several solutions to this problem, including a PMF-based enclosed chip design that passively removes the blood before readout, as well as electronic modes of detection where the blood does not interfere with the signal, which will be reported separately. The second issue is the sensitivity of the fluorescence detector. While the current design of the cell phone detector is sensitive enough for many clinical applications, we believe that optimization of the detector design will yield the necessary 10- to 20-times improvement to match the performance of table-top fluorescence scanners that are the “gold standard” for imaging microarrays. This assessment is supported by reports of mobile phone-based fluorescence imaging of individual fluorescent nanoparticles and viruses (22) and fluorescently labeled single DNA molecules (23). Third, while the ubiquitous nature of mobile phones makes it a promising platform for dedicated and nondedicated POCT platforms, there will be challenges with effectively integrating them into healthcare systems. A thorough discussion of this topic is provided in the review by Byrnes et al. (33), but in brief, issues related to developing protocols for transmitting and interpreting results and subsequently making management decisions from POC data must be resolved. Furthermore, the diversity of phone models and short life cycles in both hardware and software lead to challenges in compatibility and maintaining common standards; the slow pace of regulatory approval also introduces its own set of obstacles. However, these are challenges faced by the entire field of mobile phone-based diagnostics and are not limited to the D4 POCT. We believe the widespread dissemination of phones, their user friendliness, and their computing power make them very attractive vehicles toward democratizing access to gold-standard diagnostic testing. In conclusion, the D4 POCT, combined with the field portability of mobile phone-based imaging, may pave the way for high-performance and user-friendly diagnostic immunoassays that are virtually independent from infrastructural requirements.

Materials and Methods

SI-ATRP of POEGMA on SiO₂ Substrates. In brief, POEGMA surfaces were deposited by SI-ATRP, specifically using an activator regenerated by electron transfer approach under aqueous conditions (49, 50). The experimental conditions used to create POEGMA polymer brush coatings “grafted from” substrate surfaces for our experiment are described in extensive detail elsewhere (8). These methods involve first functionalizing the substrate surface with a bromide ATRP initiator and then immersing the substrates in polymerization solution, producing surface-tethered brushes of POEGMA ~50 nm thick.

Spotting of Antibody Microarrays. All capture/detection Ab pairs and Ag were obtained from R&D Systems. The dAbs were directly conjugated to fluorophores per the manufacturer’s instructions. The cAbs (1 mg/mL) were spotted onto POEGMA-coated substrates using a PerkinElmer Piezorray noncontact printer under ambient conditions at 1 mg/mL concentration. Spots of soluble detection reagents were composed of dAbs (1 mg/mL) mixed with excipient (1 mg/mL PEG 115,000 or 0.25 mg/mL trehalose) and printed in a similar fashion. For experiments using fingerstick blood (Fig. 2), spots of heparin were also included in arrays to maintain anticoagulation. After printing, D4 chips were placed under vacuum desiccation (30 KPa) overnight to facilitate noncovalent immobilization of Abs into the polymer brush.

D4 Immunoassay. To generate dose–response curves, D4 chips were incubated with dilution series of analyte-spiked calf serum (Clontech) or whole chicken blood (Innovative Research, Inc.) for 90 min. Substrates were then

briefly rinsed in 0.1% Tween-20/PBS then dried. Arrays were imaged on an Axon Genepix 4400 tabletop scanner (Molecular Devices, LLC). The limit-of-blank (LOB) was estimated from the mean fluorescence intensity (μ) and SD (σ) from 10 blank samples, defined as $LOB = \mu_{\text{blank}} + 1.645\sigma_{\text{blank}}$, as described by Armbruster and Pry (51). LOD was estimated from spiked low-concentration samples (LCS) above the LOB, such that $LOD = LOB + 1.645\sigma_{\text{LCS}}$. Low, intermediate, and high concentrations within the working range of each assay were used to determine the COV_{intra} and COV_{inter} . The dynamic range (DR) was defined as the range of concentrations from the LOD to the greatest concentration that had a fluorescence signal greater than 3σ of that from the next-lower concentration in the dilutions series. Data were fit to a five-parameter logistic (5-PL) fit curve (52) using OriginPro 9.0 (OriginLab Corp.). Details on the effect of shorter incubation times (15 min and 60 min) on assay results are shown in *SI Appendix, Fig. S2*.

D4 and ELISA Measurements of Patient Leptin. Approval for clinical studies was obtained from the Duke University Health System Institutional Review Board (protocol no. Pro00064838). Parental assent and subject assent were first obtained, and then 0.5–1.0 mL of blood was drawn in a lavender-top collection tube containing K2-EDTA. For D4 assays in whole blood the blood was directly applied to D4 chips immediately after collection then processed and analyzed as usual. For D4-sandwich immunoassay comparisons the tube was then centrifuged for 10 min at $1,500 \times g$, and the resulting serum was aliquoted into cryovials and stored at -80°C for later use. Samples were then thawed and analyzed with D4 and ELISA in parallel. Standard leptin sandwich immunoassays (Quantikine ELISA kit; R&D Systems) were performed per the manufacturer's instructions, which required a $100\times$ dilution with assay diluent buffer for all specimens. For D4 measurements, specimens from lean patients were directly run without dilution. In contrast, we found that two obese patient specimens had extremely high leptin levels, above the upper limit of quantitation of the leptin D4. Thus, all specimens from obese patients (to maintain consistency) were diluted $10\times$ with whole calf serum (and not sandwich immunoassay diluent buffer, to effectively maintain complexity of the biological milieu) before running the leptin D4.

Mobile-Phone Imaging of D4 Arrays. The mobile phone-based D4 assay reader device was created by integrating a 3D-printed optomechanical attachment to a smartphone (Lumia 1020; Nokia) which included an oblique illumination source for highly sensitive fluorescence detection (*SI Appendix, Fig. S5*) (22, 23). A standard microscope slide can be inserted into the smartphone at-

tachment from the side and illuminated by an excitation beam from a battery-powered laser diode (638 nm, 180 mW; Mitsubishi Electric) at an incidence of $\sim 75^\circ$ from the back side (*SI Appendix, Fig. S5A*). The fluorescence signal of the sample was collected from the other side of the glass slide by an external lens ($f_2 = 2.6$ mm; UCTronics) and passed through an emission filter (690/50 nm; Semrock) before entering the smartphone camera unit (*SI Appendix, Fig. S5A*). The smartphone camera is equipped with a 2/3-inch complementary metal-oxide semiconductor sensor with 41 megapixels and $1.12\ \mu\text{m}$ in pixel pitch. The default smartphone camera lens assembly (Carl Zeiss) has an effective focal length of $f_1 = \sim 6.86$ mm, and therefore the nominal magnification (M) of the smartphone-based fluorescence microscope is $M = \sim 2.6\times$. This mobile microscope design provides a half-pitch resolution of $\sim 0.98\ \mu\text{m}$ over an imaging field of view of $\sim 0.8\ \text{mm}^2$ as characterized earlier (53). The size of the device is about $14.5 \times 8.0 \times 7.8$ cm in length, width, and height, respectively, and the weight of the attachment excluding the smartphone is ~ 185 g. For a typical fluorescence image, an integration time of 4 s was used on the mobile phone. Fluorescence images were initially saved in raw DNG format ($7,152 \times 5,368$ pixels) on the phone memory and subsequently converted into 16-bit TIFF for further analysis. Image analysis and quantification were performed by ImageJ on a desktop computer for this study and can be conducted by a custom-developed smart application running on the same phone in future designs.

ACKNOWLEDGMENTS. We thank John Horton for technical support with microarray imaging and Dr. Michael Muehlbauer and Huaxia Cui at the Duke Molecular Physiology Institute for assistance with clinical ELISA. We also thank Dr. Robert Benjamin and Teri R. Willis for assistance with procuring clinical samples. D.Y.J. was supported by the Duke Medical Scientist Training Program (T32GM007171), and K.M. was supported by a T32 grant from the National Institute of Diabetes and Digestive and Kidney Diseases. The authors gratefully acknowledge funding support from National Heart, Lung, and Blood Institute Grants 1R21HL115410-01 and 7R41HL123871-02; National Cancer Institute Grant 1UG3CA211232-01; National Center for Advancing Translational Sciences Grant UL1TR001117; and Department of Defense United States Special Operations Command Grant W81XWH-16-C-0219. Funding support was also received from the Duke Global Health Institute, Duke-Coulter Translational Partnership, Humanitarian Innovation Fund, and European Commission. The content reflects the views of the authors and does not necessarily represent the official views of the supporting organizations.

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