analytical chemistry

¹ Direct Analysis of δ^2 H and δ^{18} O in Natural and Enriched Human Urine ² Using Laser-Based, Off-Axis Integrated Cavity Output Spectroscopy

³ Elena S. F. Berman,^{*,†} Susan L. Fortson,[†] Steven P. Snaith,[†] Manish Gupta,[†] Isabelle Chery,[‡] ⁴ Stephane Blanc,[‡] Edward L. Melanson,[§] Peter J Thomson,[⊥] and John R. Speakman^{⊥,||}

s [†]Los Gatos Research, 67 East Evelyn Ave, Suite 3, Mountain View California 94043, United States

⁶ [†]Department of Ecology, Physiology and Ethology, Hubert Curien Multidisciplinary Institute, 23 rue Becquerel, 67087 Strasbourg
⁷ cedex 2, France

8 [§]Division of Endocrinology, Metabolism, and Diabetes/Division of Geriatrics, University of Colorado Anschutz Medical Campus,

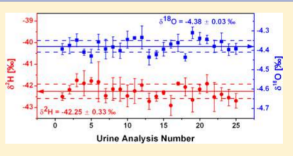
9 12801 East 17th Ave, Aurora, Colorado 80045, United States

10 ¹Institute of Biological and Environmental Sciences, University of Aberdeen, Tillydrone Ave Aberdeen, AB24 2TZ, Scotland, U.K.

11 ^{II}State Key laboratory of Molecular and Developmental Biology, Institute of Genetics and Developmental Biology, Chinese Academy

12 of Sciences, Beijing, China

ABSTRACT: The stable isotopes of hydrogen (δ^2 H) and oxygen 13 $(\delta^{18}O)$ in human urine are measured during studies of total energy 14 expenditure by the doubly labeled water method, measurement of total 15 body water, and measurement of insulin resistance by glucose disposal 16 among other applications. An ultrasensitive laser absorption spec-17 trometer based on off-axis integrated cavity output spectroscopy was 18 demonstrated for simple and inexpensive measurement of stable 19 isotopes in natural isotopic abundance and isotopically enriched human 20 urine. Preparation of urine for analysis was simple and rapid 21 (approximately 25 samples per hour), requiring no decolorizing or 22



distillation steps. Analysis schemes were demonstrated to address sample-to-sample memory while still allowing analysis of 45 natural or 30 enriched urine samples per day. The instrument was linear over a wide range of water isotopes ($\delta^2 H = -454$ to +1702 % and $\delta^{18}O = -58.3$ to +265 %). Measurements of human urine were precise to better than 0.65 % 1 σ for $\delta^2 H$ and 0.09 % 1 σ for $\delta^{18}O$ for natural urines, 1.1 % 1 σ for $\delta^2 H$ and 0.13 % 1 σ for $\delta^{18}O$ for low enriched urines, and 1.0 % 1 σ for $\delta^2 H$ and 0.08 % 1 σ for $\delta^{18}O$ for high enriched urines. Furthermore, the accuracy of the isotope measurements of human urines was verified to better than ±0.81 % in $\delta^2 H$ and ±0.13 % in $\delta^{18}O$ (average deviation) against three independent isotope-ratio mass spectrometry laboratories. The ability to immediately and inexpensively measure the stable isotopes of water in human urine is

30 expected to increase the number and variety of experiments which can be undertaken.

₃₁ Analysis of the stable isotopes of hydrogen (δ^2 H) and oxygen ₃₂ (δ^{18} O) in human body water is used in a variety of biomedical 33 applications including measurement of total energy expenditure 34 (TEE) by the doubly labeled water (DLW) method,¹⁻³ 35 measurement of total body water,⁴ and measurement of insulin 36 resistance by glucose disposal^{5,6} among other applications. 37 Currently, the vast majority of studies use isotope-ratio mass ₃₈ spectrometry (IRMS) for analysis of δ^2 H and δ^{18} O in body 39 waters. For IRMS analysis, bodily fluids (e.g., urine) require 40 either extensive purification, such as cryogenic distillation 41 followed by decolorization,⁷ or analysis by CO₂ equilibration 42 for ¹⁸O measurements and zinc or chromium reduction for ²H 43 measurements.^{8,9} These preparation methods and IRMS 44 analyses are labor-intensive, costly, and limited to only a few 45 measurement laboratories worldwide. However, in order for the 46 aforementioned biomedical applications to become widely 47 available, measurements of a large number of samples must be 48 completed quickly, accurately, and inexpensively, preferably at a 49 location near the site of sample generation.

Ultrasensitive laser absorption spectroscopy, such as off-axis 50 integrated cavity output spectroscopy (OA-ICOS) and cavity 51 ring down spectroscopy (CRDS), provides the opportunity to 52 measure δ^2 H and δ^{18} O rapidly, accurately, and inexpen-53 sively.^{10–12} Furthermore, laser-based instrumentation does 54 not require highly trained operators and has a small footprint, 55 allowing measurements to be made by researchers generating 56 the samples. While studies have shown that laser-based 57 instruments require corrections for organic contamination of 58 samples,^{11,13,14} two laboratories have recently shown that the 59 organic component of urine does not adversely affect laser-60 based isotope measurements.^{7,15} O'Grady et al. utilized CRDS 61 to measure natural isotopic abundance human urines that had 62 been either cryogenically distilled or decolorized with carbon 63 black.⁷ Thorsen et al. used CRDS to measure natural and 64

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