The Potential of Multi-Biomarker Panels in Nutrition Research: Total Fruit Intake as an Example

Aoife E. McNamara 1,2, Janette Walton 3,4, Albert Flynn 3, Anne P. Nugent 5, Breige A. McNulty 1 and Lorraine Brennan 1,2*

1 School of Agriculture and Food Science, UCD Institute of Food and Health, University College Dublin, Dublin, Ireland, 2 UCD Conway Institute of Biomolecular and Biomedical Research, University College Dublin, Dublin, Ireland, 3 School of Food and Nutritional Sciences, University College Cork, Cork, Ireland, 4 Department of Biological Sciences, Cork Institute of Technology, Cork, Ireland, 5 School of Biological Sciences, Institute for Global Food Security, Queens University Belfast, Belfast, United Kingdom

Dietary and food intake biomarkers offer the potential of improving the accuracy of dietary assessment. An extensive range of putative intake biomarkers of commonly consumed foods have been identified to date. As the field of food intake biomarkers progresses toward solving the complexities of dietary habits, combining biomarkers associated with single foods or food groups may be required. The objective of this work was to examine the ability of a multi-biomarker panel to classify individuals into categories of fruit intake. Biomarker data was measured using 1H NMR spectroscopy in two studies: (1) An intervention study where varying amounts of fruit was consumed and (2) the National Adult Nutrition Survey (NANS). Using data from an intervention study a biomarker panel (Proline betaine, Hippurate, and Xylose) was constructed from three urinary biomarker concentrations. Biomarker cut-off values for three categories of fruit intake were developed. The biomarker sum cut-offs were ≤4.766, 4.766–5.976, >5.976 µM/mOsm/kg for <100, 101–160, and >160 g fruit intake. The ability of the biomarker sum to classify individuals into categories of fruit intake was examined in the cross-sectional study (NANS) (N = 565). Examination of results in the cross-sectional study revealed excellent agreement with self-reported intake: a similar number of participants were ranked into each category of fruit intake. The work illustrates the potential of multi-biomarker panels and paves the way forward for further development in the field. The use of such panels may be key to distinguishing foods and adding specificity to the predictions of food intake.

Keywords: dietary biomarkers, nutrition, dietary assessment, food intake, metabolomics, multi-biomarker panel

INTRODUCTION

Dietary assessment is important for the elucidation of diet-disease relationships; however, traditional dietary assessment techniques are subject to some well-documented limitations (1). The identification of dietary and food intake biomarkers offers the potential of accurate and objective measure of food intake (2, 3). Food intake biomarkers are single metabolites, or a combination of metabolites, reflecting the consumption of either a specific food or food group, displaying a clear time- and dose-response after intake (4). An extensive range of putative intake biomarkers
of commonly consumed foods have been identified to date (5) but more work is needed toward confirming the utility of such biomarkers. To be efficient indicators of dietary intake, biomarkers need to be validated including analysis of sensitivity, specificity, and dose-response (6). Selecting a single intake biomarker to represent exact dietary intake is difficult due to the overlapping range of nutrients, non-nutrients, and bioactives present in foods. However, previously investigated food intake biomarkers have proven to be good predictors of consumption for classes within food groups, such as fruits and vegetables (7).

As the field of food intake biomarkers progresses, and the need to elucidate the complexities of dietary habits arises, combining biomarkers associated with single foods or food groups may be required (8). It is possible that by combining two or more biomarkers together into multi-biomarker panels could result in more sensitive and specific estimates of intake.

Development of validated and comprehensive multi-biomarker panels have the potential to add value to the assessment of dietary intake by enabling the capture of a broad range of dietary exposure including bioactive compounds, foods, food groups, and complex dietary patterns (9). Multi-biomarker panels have been previously used to classify individuals into banana consumers and non-consumers more robustly compared to individual biomarkers (10). In another example a multi biomarker panel, composed of beer ingredient and food processing biomarkers was capable of distinguishing beer consumption from urine samples collected before and up to 12h after intake of beer with excellent specificity and sensitivity (11). Other examples of successful use of multiple biomarkers include the SU.VI.MAX study, where the sum of urinary flavonoid biomarkers demonstrated higher correlations with fruit and fruit juice consumption than any of the included biomarkers individually (12). A biomarker panel containing the wine biomarkers ethyl glucuronide and tartrate outperformed individual markers when predicting wine consumers and non-consumers [90.7% receiver operating characteristics (ROC) curves area under the curve (AUC) compared to 86.3% for ethyl glucuronide and 85.7% for tartrate]. This panel was validated in an epidemiological study and was capable of estimating whether or not participants had consumed wine in the previous 3 days (13).

Multi-biomarker panels can be applied to classify and monitor adherence to dietary patterns. This was recently illustrated in post-menopausal women where a biomarker panel was capable of discriminating between high and low quintiles of adherence to four diet scores [the alternate Mediterranean diet score (aMED), alternate Healthy Eating Index (AHEI)-2010, Dietary Approaches to Stop Hypertension (DASH) diet, and the Healthy Eating Index (HEI)-2015] (14). Another interesting application of multiple biomarkers is the interrogation of the relationships between food intake and diseases incidence. In this context previous work has employed a biomarker score, derived from multiple biomarkers of fruit and vegetable intake, as a proxy for intake to examine the relationship with diabetes incidence (15). This biomarker score demonstrated an inverse association with diabetes incidence, with odds ratio (OR) of incidence decreasing with increasing intake of fruit and vegetables (highest quartile of intake compared to lowest OR: 0.13; 95% CI: 0.08, 0.21) (15). Collectively, these studies highlight the potential for combinations of multiple biomarkers in determining intake of foods or dietary patterns and assessment of relationships with health outcomes.

Fruit is an important component of a healthy diet and previous work has investigated biomarkers of various fruit. A recent systematic review of the literature conclude that we have limited knowledge for biomarkers of pome and stone fruit (16) with many biomarkers requiring validation in terms of relating to fruit intake. With respect to apple intake the most promising biomarker identified was phloretin and phloretin glucuronide. The biomarker arbutin is promising for pear intake but again requires more validation. A review of biomarkers of tropical fruit intake also highlighted a dearth of information (17), concluding clearly that there is a need for further research in the area. Proline betaine is a well-established biomarker of citrus intake with previous work indicating that urinary proline betaine concentrations can give quantitative food intake information (18, 19). However, as most consumers eat more than one fruit, there is an interest to examine total fruit intake and thus combining multiple biomarkers could be a useful approach to estimate total fruit intake.

Although previous work has indicated the potential of multiple biomarkers in the form of multi biomarker panels, combining of biomarkers to give quantitative information on food intake is not trivial. Previous studies have highlighted the potential of multi biomarker panels in terms of classification into consumers and non-consumers. However, more work is needed in the field to examine the potential of such biomarker combinations for use in assessment of consumption of different quantities of food. Therefore, the objective of this work was to examine the ability of a multi-biomarker panel to classify individuals into categories of fruit intake.

MATERIALS AND METHODS

National Adult Nutrition Survey (NANS) Study—Cross-Sectional Study

Details of the NANS study have been published elsewhere (https://www.iuna.net/) (20). Ethical approval for this study was granted by the University College Cork Clinical Research Ethics Committee of the Cork Teaching Hospitals [ECM 3 (p) 4 September 2008] and recruitment began in May 2008. Briefly, NANS collected data on habitual food and beverage consumption, lifestyle, health indicators, and attitudes to food and health in 1,500 adults, representative of the population during 2008–2010 in Republic of Ireland. A subset of this population (N = 565) was randomly selected, to ensure equal numbers of men and women across the age ranges (18–90 years) for metabolomics analysis as previously described (20). A 4 days semi weighed dietary record was used to collect dietary data over 4 consecutive days. Detailed information on the type and amount of all foods, drinks and nutritional supplements consumed over the 4 days was recorded by participants. Where possible, participants were asked to weigh foods and encouraged to retain
food packaging for further information on foods consumed, but
where weights were not recorded a photographic food atlas was
used to estimate food weights (18). Dietary data was analyzed
using WISP software (Tinuviel Software, Anglesey, UK), which
uses data from McCance and Widdowson’s The Composition of
Foods, fifth and sixth editions and all supplementary editions to
generate nutrient data (21–24). Dietary data was coded into 2552
individual food codes and grouped into one of 68 food groups.
For the purpose of this analysis any fruit containing food groups
were collapsed into a single “Total Fruit Intake” group and the
mean daily fruit intake over the 4 days was calculated. Biological
samples were collected at the end, or as close to as possible, of
the dietary recording period, including a fasting first void urine
sample in a sterile 50 mL tube which was chilled until processing.
All urine samples were centrifuged at 1,800 × g for 10 min at 4°C.
Aliquots of 1 mL were stored at −80°C for later analysis.

A-Diet Validation Intervention Study
Ethical approval for this A-Diet study was granted by the
UCD Sciences Human Research Ethics Committee (LS-17-16-
Brennan). Recruitment commenced at the end of March 2017
via advertisement (posters, flyers, and emails) and finished
in November 2017. The recruitment process is outlined in
Supplementary Figure 1. Details of the study design are
published elsewhere (25). Briefly, inclusion criteria included
healthy, non-pregnant/lactating and non-smoking individuals,
between 18 and 60 years old, and with a body mass index (BMI) between 18.5 and 30 kg/m², without any
diagnosed health condition (chronic or infectious diseases), no
consumption of medications/nutritional supplements or any
allergies/intolerances to the test foods. Once informed consent
was acquired, participants were assigned to either a lunch (N =
27) or dinner (N = 34) test meal group and invited to partake
in a 5-weeks study. Each test week participants were provided
with four portions of a test meal and asked to consume this test
meal for 4 consecutive days. During these 4 days, participants
were also asked to avoid consuming any other foods related to
the test meal ingredients. Participants were also asked to record a
4 days dietary record for each test week, to ensure compliance.
The focus of this analysis is the amount of fruit consumed as
part of these test meals. The low, medium, and high portions
of fruit provided was 50, 100, 300 g and 80, 160, 320 g for apples
and oranges, respectively.

Fasting first void urine was collected after an overnight 12 h
fast at the end of each test week and chilled until processing.
All urine samples were centrifuged at 1,800 × g for 10 min at 4°C.
Aliquots of 1 mL were stored at −80°C for later analysis.

Metabolomic Analysis of Urine Samples
Metabolomic analysis was performed using nuclear magnetic
resonance (NMR) spectroscopy. Urine samples were first
defrosted and then prepared by addition of 250 μL phosphate
buffer (0.2 mol KH₂PO₄/L, 0.8 mol K₂HPO₄/L) to 500 μL urine.
After centrifugation at 5,360 × g for 5 min at 4°C, 10 μL
sodium trimethylsilyl[2,2,3,3-²H₄] propionate (TSP) and 50 μL
deuterium oxide (D₂O) were added to 540 μL of the supernatant.
Spectra were acquired on a 600 MHZ Varian Spectrometer
(Varian Limited, Oxford, United Kingdom) by using the first
increment of a nuclear Overhauser enhancement spectroscopy
pulse sequence at 25°C. Spectra were acquired with 16,384 data
points and 128 scans. Water suppression was achieved during
the relaxation delay (2.5 s) and the mixing time (100 ms). All
¹H NMR urine spectra were referenced to TSP at 0.0 parts per
million (ppm) and processed manually with the Chenomx NMR
Suite (version 7.7) by using a line broadening of 0.2 Hz, followed
by phase and baseline correction. Three metabolites were
chosen to demonstrate the proof-of-concept that combinations
of biomarkers could be used to predict total fruit intake. The
biomarkers chosen were xylose, proline betaine, and hippurate.
Identification and quantification of metabolites was achieved
using the Chenomx library. To confirm correct assignment, a
urine sample was spiked with an analytical standard and a ¹H
NMR spectrum acquired.

Osmolality was measured by using an Advanced Micro
Osmometer model 3300 (Advanced Instruments). Aliquots of
urine were measured for osmolality with the use of micro-
osmometry by freezing point depression, with values reported as
the number of solute particles, in moles, dissolved in a kilogram
of urine (mOsm). Metabolite concentrations were normalized
to osmolality for quantifying urinary concentrations of xylose,
proline betaine and hippurate.

Statistical Analysis
Statistical analysis was performed using IBM SPSS software
package version 24.0 for windows (SPSS Inc. Chicago, IL,
USA) and SIMCA-P software (version13; Umetrics). One-way
analysis of variance was performed to compare tertiles of self-
reported intake and urinary concentrations of three biomarkers.
Spearman’s correlations were used to assess association between
mean daily self-reported total fruit intake and biomarkers. The
urinary concentrations of each biomarker were summed together
to create a single combined biomarker value for each individual.
Using the intervention data cut-offs were developed for certain
fruit intake categories.

RESULTS
Examining Relationship Between
Biomarkers and Self-Reported Fruit Intake
Urinary biomarkers of interest were quantified by ¹H NMR
analysis in the NANS cross-sectional study. The following
concentrations were obtained: xylose (range: 0.07–2.19 mM),
proline betaine (range: 0.04–2.13 mM), and hippurate (range:
0.13–15.87 mM). Participants were grouped into tertiles based
on self-reported mean daily intake of total fruit from semi-
weighed food records (Table 1, Supplementary Table 1). All
three urinary biomarker concentrations increased as the intake
of fruit increased with significant increases observed for proline
betaine and hippurate. The urinary concentrations of all three
biomarkers were also significantly correlated with mean daily
intake of fruit. The main contributors to total fruit intake were
apples, bananas, oranges, pears, strawberries, and pineapple, all
of which increased significantly across tertiles of intake except
for pineapple.
Further analysis was performed using total fruit consumers only (N = 509) (Table 2). Similar trends were observed for the urinary biomarker concentrations, increasing across tertiles of intake with significant values for proline betaine and hippurate.

Using Multiple Biomarkers to Classify Individuals Into Categories of Intake

The multi-biomarker panel was used to create a combined biomarker value by summing the individual biomarkers. In the A-Diet validation dataset (N = 160) the combined biomarker values were used to determine cut-offs for classification of individuals into categories of intake. Participants were categorized into one of three groups of intakes (0–100, 101–160, and >160 g/d) based on the fruit consumed as part of the intervention. The average sum of biomarkers for participants who consumed ≤100 g/day of fruit (4.766 μM/mOsm/kg) was set as cut-off point 1. Cut-off 3 was calculated as the average biomarker sum (5.976 μM/mOsm/kg) of participants who consumed >160 g of fruit in the A-diet study. The middle cut-off was set at any value in between cut-off 1 and 3 (Table 3).

To examine the ability of these biomarker sum values to categorize participants into categories of intake the method was applied to NANS study. Using the NANS (N = 546) self-reported dietary data participants were categorized into three groups of fruit intake (Table 4). Independently using the urinary biomarker concentrations, participants were assigned to a category of intake using the biomarker sum cut-offs. Excellent agreement between the two methods was observed with for example, 97 participants self-reporting intakes between 101 and 160 g/d, and 86 participants assigned into this category based on the biomarker panel data (Table 4 and Supplementary Table 2). Assessment of the data split by gender revealed similar agreement trends (Table 5).

In order to examine other biomarker combinations, the biomarker values that increased across categories of intake were selected (Supplementary Table 3). The ability to categorize participants into categories of intake was examined with all biomarkers and the sum of proline betaine and hippurate providing the best agreement (Table 6).

DISCUSSION

This study developed a biomarker panel that was capable of classifying individuals into categories of fruit intake. Examination of the approach in a free-living cross-sectional study revealed excellent agreement with self-reported intake. Similar number of participants were ranked into each category of fruit intake. The work illustrates the potential of multi-biomarker panels and paves the way forward for further development in the field.

The three biomarkers chosen to build the multi-metabolite panel in this research have frequently been found in previous research to be associated with fruit intake. Xylose was previously identified as a food intake biomarker associated with apple intake and capable of ranking participants in an observational study by increasing intake (25). Proline betaine is a well-established, robust, and quantitative biomarker of citrus intake (18, 26). Proline betaine was previously included as part of a biomarker panel for the investigation of orange juice intake and juice quality (27). Hippurate has often been associated with consumption of polyphenol rich plant foods, such as citrus fruit, bananas, and berries (10, 26, 28–30). This research combined these three food intake biomarkers into a biomarker score for the successful ranking of fruit intake. It is important to acknowledge that there are other potential biomarkers of fruit intake; however, examination of the full panel of potential biomarkers was beyond the scope of the current work where the emphasis was on the demonstration of the potential of combination of biomarkers.

Previous multi-metabolite panels have focused on associating the panels with intake of certain foods or achieving a dichotomous classification of consumers and non-consumers. An example of previous work demonstrated good associations

| Table 1 | Urinary food intake biomarker concentrations across tertiles of fruit intake in cross-sectional study (NANS). |
| --- | --- | --- | --- | --- | --- | --- |
| Metabolite (µ/ M/mOsm/kg) | Tertile 1 (N = 180) | Tertile 2 (N = 181) | Tertile 3 (N = 185) | Mean daily intakes (g/d) | Hippurate | Proline betaine | Hippurate |
| Mean | SD | Mean | SD | Mean | SD | p-value | Rho | p-value |
| Metabolite (µ/ M/mOsm/kg) | Xylose | 0.596 | 0.34 | 0.614 | 0.35 | 0.637 | 0.42 | 0.074 | 0.078 |
| | Proline betaine | 0.285 | 0.22 | 0.415 | 0.34 | 0.569 | 0.44 | <0.001 | 0.400 |
| | Hippurate | 3.600 | 2.78 | 3.991 | 2.87 | 4.727 | 3.20 | 0.001 | 0.181 |
| Mean daily intakes (g/d) | Total fruit | 28.42 | 24.61 | 127.80 | 32.61 | 315.19 | 115.44 | <0.001 |
| | Apples | 4.94 | 13.50 | 23.93 | 30.76 | 58.15 | 65.78 | <0.001 |
| | Banana | 7.17 | 15.28 | 26.86 | 31.77 | 43.07 | 42.70 | <0.001 |
| | Oranges | 2.49 | 9.86 | 29.56 | 42.55 | 94.14 | 105.97 | <0.001 |
| | Pears | 0.52 | 3.70 | 4.46 | 15.61 | 22.18 | 47.46 | <0.001 |
| | Strawberry | 1.03 | 5.41 | 2.48 | 9.34 | 4.44 | 13.40 | 0.005 |
| | Pineapple | 0.47 | 3.73 | 1.20 | 7.47 | 2.75 | 16.91 | 0.129 |

All values presented are mean and standard deviation (SD) unless stated otherwise. ANOVA, analysis of variance; MDI, Mean daily intake. N = 546.

* Spearman’s correlation between each biomarker and total mean daily fruit intake.
between biomarkers of fruit and vegetables intake and self-reported dietary records (12). When 24 h urinary concentrations of isorhamnetin, hesperetin, naringenin, kaempferol, and phloretin were combined in a panel they were correlated with fruit intake \( (r = 0.27, p = 0.06) \), juice fruit intake \( (r = 0.28, p = 0.04) \), and intake of total fruits and fruit juices \( (r = 0.38, p = 0.006) \). The correlations for the fruit related panel improved when examined in spot urine samples: fruit intake \( (r = 0.34, p = 0.01) \), juice fruit intake \( (r = 0.44, p = 0.001) \), and total fruit and juice intake \( (r = 0.47, p = 0.0004) \). The authors concluded that this combination of flavonoids could be used as a reliable biomarker of total fruit and juice intake, however, to the best of our knowledge there is no demonstration that the biomarkers could predict intake. Our work is an important advancement as it clearly demonstrates that the multi-biomarker approach is capable of a classification of intake into a range of categories. However, it should be noted that the present work did not make the distinction between whole fruit and fruit juices and the biomarker panel classification was based on total fruit intake including juices.

Our previous work identified four food intake biomarkers of sugar sweetened beverages (formate, citrulline, taurine, and isocitrate) using heat-map analysis of metabolomic urinary profiles from the NANS study (31). These markers were confirmed by food analysis of the sugar-sweetened beverage and an acute intervention study. The markers were combined in a panel and ROC curves demonstrated that the panel could discriminate between consumers and non-consumers of sugar-sweetened beverages \( (AUC = 0.8) \) and was more

| TABLE 2 | Urinary food intake biomarker concentrations across tertiles of fruit intake in cross-sectional study (NANS), total fruit consumers only. |
|---|---|---|---|---|---|---|---|---|
| Fruit intake category (g/d) | Tertile 1 \( (N = 168) \) | Tertile 2 \( (N = 168) \) | Tertile 3 \( (N = 173) \) | One-way ANOVA | Spearman’s correlation |
| | Mean | SD | Mean | SD | Mean | SD | p-value | Rho | p-value |
| Metabolite \( (\mu M/mOsm/kg) \) | | | | | | | | | |
| Xyllose | 0.601 | 0.35 | 0.604 | 0.35 | 0.649 | 0.43 | <0.001 | 0.424 | 0.090 | 0.043 |
| Proline betaine | 0.298 | 0.24 | 0.425 | 0.33 | 0.581 | 0.45 | <0.001 | 0.382 | <0.001 |
| Hippurate | 3.622 | 2.63 | 3.957 | 2.82 | 4.771 | 3.24 | 0.001 | 0.187 | <0.001 |
| Mean daily intakes (g/d) | | | | | | | | | |
| Total fruit | 42.67 | 26.44 | 139.58 | 32.28 | 323.36 | 114.98 | <0.001 |
| Apples | 8.66 | 18.54 | 25.60 | 33.86 | 59.09 | 66.35 | <0.001 |
| Banana | 10.05 | 17.90 | 29.61 | 32.63 | 43.12 | 43.46 | <0.001 |
| Oranges | 4.18 | 13.92 | 33.34 | 43.80 | 97.75 | 108.18 | <0.001 |
| Pears | 1.11 | 6.39 | 4.24 | 15.46 | 23.71 | 48.71 | <0.001 |
| Strawberry | 1.36 | 6.01 | 2.42 | 9.45 | 4.74 | 13.81 | 0.008 |
| Pineapple | 1.07 | 7.08 | 1.14 | 7.21 | 2.55 | 16.75 | 0.399 |

All values presented are mean and standard deviation (SD) unless stated otherwise. ANOVA, analysis of variance; MDI, Mean daily intake. N = 509.

* Spearman’s correlation between each biomarker and total mean daily fruit intake.

| TABLE 3 | Cut-off points for each of the fruit intake categories derived from the Intervention Study. |
|---|---|
| Fruit intake category (g/d) | Biomarker cut-offs \( (\mu M/mOsm/kg) \) |
| 0–100 | <4.766 |
| 101–160 | 4.766–5.974 |
| >160 | ≥5.975 |

Cut-offs were calculated using sums of biomarker concentrations in each intake category from the A-Diet intervention study.

| TABLE 4 | Classification into categories of fruit intake based on biomarker data or based on self-reported intake data in the cross-sectional study (NANS). |
|---|---|---|
| Fruit intake category (g/d) | Self-reported data (N) | Biomarker data (N) |
| 0–100 | 227 | 306 |
| 101–160 | 97 | 88 |
| >160 | 222 | 154 |

Participants were classified into one of 3 categories based on either (1) Self-reported dietary data or (2) biomarker data. The number of participants into each category is reported, N = 546.

| TABLE 5 | Distribution of participants classified into each category of intake based on sum of urinary biomarker concentrations compared to reported intake data in the cross-sectional study, split by gender. |
|---|---|---|---|---|---|---|
| NANS | Self-reported | Predicted |
| Gender | M | F | M | F |
| Intake category | | | | | | |
| 0–100 (g/d) | 128 | 99 | 170 | 136 |
| 101–160 (g/d) | 42 | 55 | 42 | 44 |
| >160 (g/d) | 108 | 114 | 66 | 88 |

Participants \( (N = 546) \) categories based on cut-offs were calculated using sums of biomarker concentrations in each intake category from the A-Diet intervention study; ADV, A-Diet Validation Study; NANS, National Adult Nutrition Survey; M, Male; F, Female; N, Number of participants.
predictive of intake than the individual biomarkers themselves (AUCs ranging from 0.5 to 0.7). A recently published study used data from the KarMeN study to identify five metabolites [methoxyeugenol glucuronide (MEUG-GLUC), dopamine sulfate (DOP-S), salsolinol sulfate, 6-hydroxy-1-methyl-1,2,3,4-tetrahydra-β-carboline sulfate, and xanthurenic acid] that were discriminative between high and non-consumers of banana (10). Individually, DOP-S had the best prediction ability [AUC = 0.84, error rate (ER) = 0.25] for classifying high consumers against non-consumers but was not as robust as a combination of all five metabolites (AUC = 0.90, ER = 0.13). However, the best predictive ability was a combined panel of MEUG-GLUC and DOP-S with the lowest error rate of misclassification. This research demonstrates how a panel of food intake biomarkers which individually were not robust enough, when combined can be used to classify recent banana intake. Our research takes the application of biomarker panels a step further by classifying participants into categories of fruit consumed and moves beyond the dichotomous classification of consumer/non-consumer.

While the above work demonstrates the potential for multi-biomarkers in terms of estimating food intake such panels can also be used to assess the relationship between food and diet-related diseases. A combined biomarker-score was developed using the standardized plasma values of vitamin C, β-carotene, and lutein, all of which were previously related to fruit and vegetable intake (15). This score was inversely associated with odds of incidence of type 2 diabetes (OR: 0.13; 95% CI: 0.08, 0.21) even after adjustment for lifestyle factors and demographics. An identified plasma biomarker panel representative of dietary habits consisted of β-alanine (beef intake), alkylresorcinols (wholegrain/rye), eicosapentaenoic acid and 3-carboxy-4-methyl-5-propyl-2-furanpropanoic acid (fish), lauric acid (saturated fats), linoleic acid (seeds, nuts, and vegetable oils), oleic acid (olive and rapeseed oil), and α and γ tocopherol (32). This combined panel was capable of predicting new cases of type 2 diabetes over a 5-years follow-up period with a specificity and sensitivity similar to classic diabetes predictors (serum adiponectin, insulin resistance, impaired glucose tolerance and impaired fasting glycaemia). Collectively, these and other studies highlight the potential of a multi-metabolite panel for the assessment of the relationship between diet and health/disease.

If future studies develop comprehensive and validated multi-biomarker panels, they could add value to the assessment of dietary intake by enabling the capture of a broad range of dietary exposures including bioactive compounds, foods, food groups, and complex dietary patterns. Panels could then be used in epidemiological research to elucidate the mechanisms and metabolic pathways of diet-related diseases and to validate self-reported dietary data. Further development of more comprehensive panels could enable measurement of adherence to specific dietary patterns, such as the Mediterranean diet. Future challenges for the field will be finding the simplest combination of metabolites to accurately determine exposures as well as validating these panels to a standard where they can be applied in nutritional research and public health surveys (9). As the field develops further, there will be a need to develop new statistical tools to integrate multiple biomarkers with self-reported data. Our work has recently developed calibration equations based on biomarker-predicted citrus intakes to gain a more accurate and objective measure of true intake (33). Using biomarker data to correct self-reported data for food intake is a promising option and could be adapted to include a biomarker panel. Further work is to develop the statistical tools to achieve this.

This study has strengths and limitations. A limitation worth noting is the fact that we examined the ability of the biomarker panel to categorize fruit intake in a cross-sectional study where intake was estimated with self-reported data. Future studies where comparison is performed in large intervention studies would be useful to examine relationships with true food intake. On the other hand, including the development of a food intake biomarker panel with testing in cross-sectional study is a strength as it demonstrates that the panel was capable of ranking fruit intake at a population level, against the background of exposure to various other foods. This research did not examine the potential impact of gut microbiome on the biomarker panel. Future work is warranted to address this.

To conclude, this study successfully demonstrated the utility of a panel of biomarkers for estimating fruit intake. The identification of comprehensive and validated multi-biomarker panels related to certain foods will be important as this field develops. The use of such panels may be key to distinguishing foods and adding specificity to the predictions of food intake. Combining such panels with self-reported measures will be important for increasing the accuracy of dietary assessment methods. Furthermore, there is potential for the use of such panels in large epidemiological studies to examine the relationships between diet and disease.

### DATA AVAILABILITY STATEMENT

The original contributions presented in this study are included in the article/Supplementary Material,
further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by UCD HREC. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

AM conducted the A-Diet Validation study, acquired, analyzed the data, and prepared the manuscript. LB designed the research, analyzed the data, and prepared the manuscript. BM, AN, JW, and AF contributed the data from the NANS study. All authors read and accepted the final version of the manuscript.

REFERENCES

1. Bingham SA. Biomarkers in nutritional epidemiology. Public Health Nutr. (2002) 5:821–7. doi: 10.1079/PHN2002368
2. Gibbons H, O’Gorman A, Brennan L. Metabolomics as a tool in nutritional research. Curr Opin Lipidol. (2015) 26:30–4. doi: 10.1097/00071145-201501000-000140
3. Jenab M, Slimani N, Bictash M, Ferrari P, Bingham SA. Biomarkers in nutritional epidemiology: applications, needs and new horizons. Hum Genet. (2009) 125:507–25. doi: 10.1007/s00439-009-0662-5
4. Gao Q, Praticò G, Scalbert A, Vergères G, Kolehmainen M, Manach C, et al. A scheme for a flexible classification of dietary and health biomarkers. Genes Nutr. (2017) 12:1–15. doi: 10.1186/s12263-017-0587-x
5. Lloyd AJ, Willis ND, Wilson T, Zubair H, Chambers E, García-Perez I, et al. Addressing the pitfalls when designing intervention studies to discover and validate biomarkers of habitual dietary intake. Metabolomics. (2019) 15:72. doi: 10.1007/s11306-019-1532-3
6. Dragsted LO, Gao Q, Scalbert A, Vergères G, Kolehmainen M, Manach C, et al. Validation of biomarkers of food intake-critical assessment of candidate biomarkers. Genes Nutr. (2018) 13:14. doi: 10.1186/s12263-018-0603-9
7. Baldrick FR, Woodside JV, Elborn JS, Young IS, McKinley MC. Biomarkers of fruit and vegetable intake in human intervention studies: a systematic review. Crit Rev Food Sci Nutr. (2011) 51:795–815. doi: 10.1080/10408398.2010.482217
8. Brennan L, Hu FB. Metabolomics-based dietary biomarkers in nutritional epidemiology-current status and future opportunities. Mol Nutr Food Res. (2019) 63: e1701064. doi: 10.1002/mnfr.201701064
9. García-Aloy M, Rabassa M, Casas-Agustench P, Hidalgo-Liberona N, Llorach R, Andres-Lacueva C. Novel strategies for improving dietary exposure assessment: Multiple-data fusion is a more accurate measure than the traditional single-biomarker approach. Trends Food Sci Technol. (2017) 69:220–9. doi: 10.1016/j.tifs.2017.04.013
10. Vázquez-Manjarrez N, Weinert CH, Ulaszewska MM, Mack CI, Micheau P, Pétra M, et al. Discovery and validation of banana intake biomarkers using untargeted metabolomics in human intervention and cross-sectional studies. J Nutr. (2019) 149:1701–13. doi: 10.1093/jn/nxz125
11. Giurdeniz G, Jensen MG, Meier S, Bech L, Lund E, Dragsted LO. Detecting beer intake by unique metabolite patterns. J Proteome Res. (2016) 15:4544–56. doi: 10.1021/acs.jproteome.6b00635
12. Mennenn LI, Sapinho D, Ito H, Bertrais S, Galan P, Hercberg S, et al. Urinary flavonoids and phenolic acids as biomarkers of intake for polyphenol-rich foods. Br J Nutr. (2006) 96:191–8. doi: 10.1079/BJN20061808
13. Vázquez-Fresno R, Llorach R, Urpi-Sarda M, Khyemenets O, Bulló M, Corella D, et al. An NMR metabolomics approach reveals a combined-biomarkers model in a wine intervention trial with validation in free-living individuals of the PREDIMED study. Metabolomics. (2015) 11:797–806. doi: 10.1007/s11306-014-0735-x
14. McCullough ML, Malniak ML, Stevens VL, Carter BD, Hodge RA, Wang Y. Metabolomic markers of healthy dietary patterns in US postmenopausal women. Am J Clin Nutr. (2019) 109:1439–51. doi: 10.1093/ajcn/nqz385
15. Cooper AJ, Sharp SJ, Luben RN, Khaw KT, Wareham NJ, Forouhi NG. The association between a biomarker score for fruit and vegetable intake and incident type 2 diabetes: the EPIC-Norfolk study. Eur J Clin Nutr. (2015) 69:449–54. doi: 10.1038/ejcn.2014.246
16. Ulaszewska M, Vázquez-Manjarrez N, García-Aloy M, Llorach R, Mattivi F, Dragsted LO, et al. Food intake biomarkers for apple, pear, and stone fruit. Genes Nutr. (2018) 13:29. doi: 10.1186/s12263-018-0620-8
17. Vázquez-Manjarrez N, Ulaszewska M, García-Aloy M, Mattivi F, Pratico G, Dragsted LO, et al. Biomarkers of intake for tropical fruits. Genes Nutr. (2020) 15:11. doi: 10.1186/s12263-020-00670-4
18. Gibbons H, Michielsen CJR, Rundle M, Frost G, McNulty BA, Nugent AP, et al. Demonstration of the utility of biomarkers for dietary intake assessment; proline betaine as an example. Mol Nutr Food Res. (2017) 61:1700037. doi: 10.1002/mnfr.201700037
19. Mack CI, Weinert CH, Egert B, Ferrario PG, Bab A, Hoffmann I, et al. The complex human urinary sugar profile: determinants revealed in the cross-sectional KarMeN study. Am J Clin Nutr. (2018) 108:502–16. doi: 10.1093/ajcn/nqz313
20. Cashman KD, Muldowney S, McNulty B, Nugent A, FitzGerald AP, Kiely M, et al. Vitamin D status of Irish adults: findings from the National Adult Nutrition Survey. Br J Nutr. (2013) 109:1248–56. doi: 10.1017/S0007114512003212
21. Nelson M, Atkinson M, Meyer J. A Photographic Atlas of Food Portion Sizes. London: Ministry of Agriculture, Fisheries and Food (2002).
22. McCance R, Widdowson E. McCance and Widdowson’s The Composition of Foods. 5th ed. Cambridge: Royal Society of Chemistry (1991).
23. McCance R, Widdowson E. McCance and Widdowson’s The Composition of Foods. 6th ed. Cambridge: Royal Society of Chemistry (2002).
24. McCance R, Widdowson E. McCance and Widdowson’s The Composition of Foods. 7th ed. Cambridge: Royal Society of Chemistry (2014).
25. McNamara AE, Collins C, Harsha PSCS, González-Peña D, Gibbons H, McNulty BA, et al. Metabolomic-based approach to identify biomarkers of apple intake. Mol Nutr Food Res. (2020) 64:e1901158. doi: 10.1002/mnfr.201901158
26. Heinzmman SS, Brown JJ, Chan Q, Bictash M, Dumas ME, Kochhar S, et al. Metabolic profiling strategy for discovery of nutritional biomarkers: proline betaine as a marker of citrus consumption. Am J Clin Nutr. (2010) 92:436–43. doi: 10.3945/ajcn.2010.29872

FUNDING

This work was supported by a H2020 European Research Council (647783).

ACKNOWLEDGMENTS

The authors would like to thank all volunteers for their commitment and patience during the study. The authors would like to acknowledge the help of Dr. Helena Gibbons and Dr. Diana González-Peña.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fnut.2020.577720/full#supplementary-material
27. Rangel-Huerta OD, Aguillera CM, Perez-de-la-Cruz A, Vallejo F, Tomas-Barberan F, Gil A, et al. A serum metabolomics-driven approach predicts orange juice consumption and its impact on oxidative stress and inflammation in subjects from the BIONAOS study. Mol Nutr Food Res. (2017) 61:1600120. doi: 10.1002/mnfr.201600120

28. Toromanović J, Kovac-Besović E, Sapcanin A, Tahirović I, Rimpapa Z, Kroyer G, et al. Urinary hippuric acid after ingestion of edible fruits. Bosn J Basic Med Sci. (2008) 8:38–43. doi: 10.17305/bjbms.2008.2994

29. Andersen MB, Kristensen M, Manach C, Pujos-Guillot E, Poulsen SK, Larsen TM, et al. Discovery and validation of urinary exposure markers for different plant foods by untargeted metabolomics. Anal Bioanal Chem. (2014) 406:1829–44. doi: 10.1007/s00216-013-7498-5

30. Walsh MC, Brennan L, Pujos-Guillot E, Sébédio JL, Scalbert A, Fagan A, et al. Influence of acute phytochemical intake on human urinary metabolomic profiles. Am J Clin Nutr. (2007) 86:1687–93. doi: 10.1093/ajcn/86.5.1687

31. Gibbons H, McNulty BA, Nugent AP, Walton J, Flynn A, Gibney MJ, et al. A metabolomics approach to the identification of biomarkers of sugar-sweetened beverage intake. Am J Clin Nutr. (2015) 101:471–7. doi: 10.3945/ajcn.114.095604

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 McNamara, Walton, Flynn, Nugent, McNulty and Brennan. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.