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Quality assurance and quality control processes

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DOI: 10.1007/s11306-017-1188-9

Document Version Peer reviewed version

Citation for published version (Harvard):

Dunn, WB, Broadhurst, DI, Edison, A, Guillou, C, Viant, MR, Bearden, DW & Beger, RD 2017, 'Quality assurance and quality control processes: summary of a metabolomics community questionnaire', Metabolomics, vol. 13, no. 5, 50. https://doi.org/10.1007/s11306-017-1188-9

Link to publication on Research at Birmingham portal

Publisher Rights Statement:

Checked for eligibility: 24/05/2017. The final publication is available at Springer via http://dx.doi.org/10.1007/s11306-017-1188-9.

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1 2	2	metabolomics community questionnaire
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45 46	25	Keywords: metabolomics, quality assurance, quality control, questionnaire,
47 48	26	Metabolomics Society
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28 Abstract

30 Introduction

The Metabolomics Society Data Quality Task Group (DQTG) developed a questionnaire regarding quality assurance (QA) and quality control (QC) to provide baseline information about current QA and QC practices applied in the international metabolomics community.

Objectives

The DQTG has a long-term goal of promoting robust QA and QC in the
metabolomics community through increased awareness via communication,
outreach and education, and through the promotion of best working practices.
An assessment of current QA and QC practices will serve as a foundation for
future activities and development of appropriate guidelines.

43 Method

QA was defined as the set of procedures that are performed in advance of
analysis of samples and that are used to improve data quality. QC was defined as
the set of activities that a laboratory does during or immediately after analysis
that are applied to demonstrate the quality of project data. A questionnaire was
developed that included 70 questions covering demographic information, QA
approaches and QC approaches and allowed all respondents to answer a subset
or all of the questions.

52 Result

The DQTG questionnaire received 97 individual responses from 84 institutions
in all fields of metabolomics covering NMR, LC-MS, GC-MS, and other analytical
technologies.

57 Conclusion

There was a vast range of responses concerning the use of QA and QC approaches that indicated the limited availability of suitable training, lack of Standard Operating Procedures (SOPs) to review and make decisions on quality, and limited use of standard reference materials (SRMs) as OC materials. The DQTG QA/QC questionnaire has for the first time demonstrated that QA and QC usage is not uniform across metabolomics laboratories. Here we present recommendations on how to address the issues concerning QA and QC measurements and reporting in metabolomics.

Introduction

Metabolomics is a scientific approach applied to the systems analysis of metabolism [Dunn 2011] operating in microbes, plants and animals [Furusawa 2013; Kusano 2015; Cheng 2015]. The discipline of metabolomics is less than 20 years of age [Oliver 1998] although the roots are much older [Pauling 1971]. Metabolomics studies typically use a pipeline from experimental design through analytical measurements (sample preparation and data acquisition) to bioinformatics processing (data processing and statistical analysis) [Brown 2005]. The validity of and confidence in the biological conclusions resulting from this pipeline are highly dependent on the quality of the procedures applied during the metabolomics study. The appropriate application of quality assurance (QA) and quality control (QC) processes are important but are often overlooked in metabolomics. In targeted metabolite studies, guidelines are available to guide the scientist in some aspects of the process including the most frequently applied Food and Drug Administration (FDA) guidelines for bioanalytical method validation [http://www.fda.gov/downloads/Drugs/Guidance/ucm070107.pdf.] as well as other materials [Garfield 2000; Hibbert 2007; Westgard 2008; Booth 2015]. However, there are currently no clear guidelines for untargeted metabolomic studies.

The Metabolomics Society's mission includes 'To promote the growth and development of the field of metabolomics internationally" [Metabolomics Society website]. To address this mission, several scientific task groups have been established to act for the community in areas requiring international community consensus. One of these is the Data Quality Task Group (DQTG) chaired by Drs. Daniel Bearden and Richard Beger. The DQTG promotes robust QA and QC in the metabolomics community through increased awareness via communication, outreach and education, and through the promotion of best working practices [Bearden 2014; Metabolomics Society task group website]. One objective of this task group is to define the current application levels of QA and QC processes in both targeted and untargeted studies across all applications in metabolomics. To complete this objective, the task group operated a questionnaire for 2 months (August September 2015) via the SurveyMonkey website (https://www.surveymonkey.com), which was advertised via e-mail alerts, Metabolomics Society web pages, Twitter and MetaboNews newsletters. The questionnaire included 70 questions covering demographic information. OA approaches and QC approaches and allowed all respondents to answer a subset or all of the questions. All responses are available in the supplementary information and on the Metabolomics Society website [13]. Here we will summarize the most important information and facts derived from the questionnaire and a number of important recommendations.

The respondents

- 97 respondents
- 36 % were principal investigators (PIs) or group leaders, 14 % were staff scientists, 20 % were post-doctoral researchers and 19 % were PhD students.

	115	• 11 % of respondents had less than 2 years of experience in metabolomics
1	116	with 31 % having greater than 8 years experience.
2	117	• The respondents applied metabolomics in a diversity of different
4	118	applications and many respondents worked across multiple disciplines
5	119	including clinical sciences (65 %), toxicology (35 %) and systems biology
6	120	(45 %).
7 0	121	• 70 % responded as working in a combination of a biological/chemical
8 9	122	laboratory and data processing/bioinformatics.
10	123	• Greater than 70 % of respondents worked with cells, biofluids and tissues
11	124	and investigated microbes (42%), plant (34%), mammals (62%) and
12	125	humans (76%). 73% and 88% of respondents applied targeted and
14	126	untargeted assays, respectively, with 34 % applying NMR spectroscopy in
15	127	their studies, 83 % applying liquid chromatography-mass spectrometry
16	128	and 50 % applying gas chromatography-mass spectrometry.
17 10	129	• 74% of respondents investigated less than 200 samples in a typical
19	130	biological study and 63 % studied less than 5000 total biological samples
20	131	each year.
21	132	
22	133	Training
23 24	134	Quality processes include training and competence assessment to ensure a
25	135	minimum quality-level is associated with processes involving staff. 65 % of 94
26	136	responses defined that they operate in an environment with no in-house training
27	137	program and 74% were not required to be involved in ongoing continuous
20 29	138	professional education. In environments where training was conducted (33
30	139	responses), professional staff (49%) and post-doctoral/graduate staff (36%)
31	140	were the major providers of training. Where training is provided, only 21 % of
32	141	instrument operators have to pass a certification test after training, with 57 %
33 34	142	applying professional staff to perform the assessment. 79 % of 85 responses do
35	143	not operate in an environment where there was a requirement to pass a
36	144	certification test after training. 73 % of 33 responses applied periodic checks of
37	145	professional practice with 58% of checks performed by professional staff as
38 39	146	indicated by 33 responses.
~ ~		

Standard Operating Procedures

The mistakes that can be introduced into metabolomics experiments through improper or inconsistent pre-analytical or analytical procedures may cause the data to be inaccurate or invalid, and this may lead to erroneous findings and conclusions. For examples see [Gika 2008; Bernini 2011; Kamlage 2014; Dunn 2012]. Consistent procedures as simple as pipetting, balance usage, sample cross-contamination control, proper preparation of solvents and sample extraction techniques all contribute to the veracity of the analytical measurements and should be thoroughly documented in Standard Operating Procedures (SOPs) and enforced in training programs. For long-term studies or interlaboratory studies, SOPs are essential for communicating well and ensuring the consistency of the data.

Eighty-seven respondents answered questions related to SOPs. SOPs were available in the laboratories of 76 % of respondents with 58 % developed in-house and a further 37 % developed from in-house and published methods.

When investigated in more detail, 90 % of respondents had access to SOPs for sample extraction, 53 % for sample storage, 75 % for analytical instruments, 52 % for assessment of data from QC samples and 33 % for deciding when QC data from instrumental analysis has failed and defining how to correct the instrumental data. As a matter of concern and shown for 84 responses was that б 70 % of respondents did not have access to a protocol for independent review of quality-related results (Figure 1A) and 80 % did not have access to a written protocol of QA review criteria (Figure 1B).

Sample measurement validation

The majority of respondents (82 responses) validate sample measurements with 73 % using repeat sample extractions and analyses, 87 % performing repeated analysis of the same sample and 54 % analyzing a historical sample periodically (Figure 2). 88% of 80 responses analyze a blank sample with extraction performed as for biological samples. Blank samples were analyzed either at the start and end of the analytical batch (28%), at regular intervals (44%) or randomly (21%) as defined in 68 responses. 78% of respondents operated a process to reduce carryover (80 responses) and 91 % randomize the order of sample analysis (80 responses). 94 % operated instrument condition checks and 79 % of 80 responses did not apply standard reference materials (SRMs); when applicable, 47 % applied a SRM once or less than once a day and 16 % greater than once per day. Methods for reporting of QC data were variable in the 80 responses collected; 34 % reported precision measurements for each metabolite, 45 % report a single range of precisions for all metabolites, 24 % report QC data on a boxplot, 56 % visualize QC samples on a PCA scores plot and 56 % provide a descriptive statement of the QC results.

QC samples

Of 80 responses, 83 % of respondents applied pooled project materials and 48 % applied standard reference materials (SRMs) as OC materials. This contradicts the results for SRM use as defined above in the sample measurement validation section. Figure 3 illustrates how often QC samples were applied for different processes including the assessment of consistency in sample preparation (80 %) and chromatography column integrity (76%). Importantly, 59% of respondents applied replicate extractions and 69% applied replicate analytical measurements with 85 % analyzing individual samples and 15 % analyzing a single pooled sample.

Data storage

Of 84 responses, 89 % store data in an archive, with 95 % of data storage being performed in an in-house archive. A lower percentage (73%) archived QA/QC data.

Inter-laboratory comparisons

Laboratory accreditation

Of 82 responses, 33 % had participated in an inter-laboratory comparison study and 48% were interested in participating in a future inter-laboratory comparison.

	213	Of 85 responses, 89 % were not required to meet laboratory accreditation and
1	214	74 % were not voluntarily attempting to meet any accreditation.
2	215	· · · / · · · · · · · · · · · · · · · ·
3	215	
4	210	
5	217	
6	218	Biggest issues in QA and QC implementation and processes
7	219	The most frequent comments related to the currently regarded biggest issues in
8	220	OA and OC are detailed below:
10	221	 Training including staff turnover and lack of training available outside the
11	221	• I failing including stall turnover and lack of training available outside the
12	ZZZ	organization
13	223	• SOP formalization, consistency and maintenance including reported
14	224	changes to published methods (for example papers published in <i>Nature</i>
15	225	Protocols
16	226	 Ensuring routing compliance to SOPs and OA processos
17	220	• Ensuring routine compliance to sol s and QA processes
18	227	• Insufficient control over sample collection and sampling consistency
19	228	• Inadequate availability of reference standards, isotopically labeled
20	229	compounds, QC samples and SRMs
21	230	 Providing a balance between OA/OC and sample throughput
22	221	 Of does not contribute to assessment of output by the wider community.
23	201	• QC does not contribute to assessment of output by the while community
24	232	and there is a need for true standards across the community
25 26	233	 A global strategy for QA/QC and its review is required
20 27	234	• Establishment of QC acceptance criteria as currently there is a lack of
28	235	reported OC results and acceptance criteria
29	236	 Additional measures beyond pooled OC samples
30	230	• Additional measures beyond pooled de samples
31	237	
32	238	Key conclusions and recommendations
33	239	1. The level of training, both in-house and external to the organization, is low;
34	240	65 % of responses replied that they operate in an environment with no in-house
35	241	training program. 74 % of responses were not required to be involved in ongoing
36	242	continuous professional education.
3/	243	Recommendation: Enhance training focused on OA and OC available as online and
30 20	213	face to face courses (for example, the Dirmingham Matchelomics Training Control
40	244	Juce-to-Juce courses (for example, the Dirininghum Metabolomics Training Centre
41	245	operates a 2-day course focused on QA and QC processes).
42	246	
43	247	2. 76 % of respondents applied SOPs. However, 70 % of respondents did not
44	248	have access to a protocol for review of quality and 80 % did not have access to
45	249	protocols focused on a review of quality processes.
46	250	Pacommondation: Appropriate gaancies and the Metabolomics Society should
47	250	Recommendation. Appropriate agencies and the metabolomics society should
48	251	provide guidance on quality assurance processes and their review; develop
49	252	consensus processes through specialist meetings and reports.
50	253	
51	254	3. The majority of respondents validate sample measurements, apply sample
52 53	255	blanks, apply protocols to minimize sample carryover and randomize the
54	256	analysis order of hiological samples
55	250	Decommondation. To provide education to the metabolomics community with an
56	257	Recommendation. To provide education to the metabolomics community, with an
57	258	emphasis on early career scientists, on sample measurement validation, and to
58	259	provide continuing education to ensure these good practices continue.
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- 4. 83 % of respondents applied pooled project materials and 48 % applied standard reference materials (SRMs) as QC materials. 59% of respondents applied replicate extractions and 69% applied replicate analytical measurements. Recommendation: To provide education to the metabolomics community, with an
- б emphasis on early career scientists, on usage of quality materials, and to provide continuing education to ensure these good practices continue.

- 5.79 % of respondents did not access SRM materials.
- Recommendation: To communicate with the metabolomics community to define the types and volumes of SRMs required.
- 6. 33 % had participated in an inter-laboratory comparison study and 48% were interested in participating in a future inter-laboratory comparison.
- Recommendation: To communicate with the metabolomics community to define the types and frequency of inter-laboratory comparison exercises and encourage independent agencies to support inter-laboratory exercises.
- 7.89% of respondents were not required to meet laboratory accreditation and 74 % were not voluntarily attempting to meet any accreditation.
- Recommendation: To investigate the requirement for laboratory accreditation with the regulatory agencies, funding bodies, the Metabolomics Society and the metabolomics community.
- 8. There is little incentive for laboratories to improve their QA/QC practices, especially given the non-trivial costs associated with a thorough QA/QC program.
- Recommendation: Recognizing the need to provide further incentive for laboratories to improve overall QA/QC practices, expert panels should be convened to develop workable, practical QA/QC recommendations and guidelines. One possible mechanism is a workshop currently being planned for later in 2017 that will define appropriate QA/QC frameworks that may be adopted widely in laboratories and, possibly, by funders, data repositories and scientific publishers.

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Compliance with Ethical Standards

The authors have defined that there are no potential conflicts of interest. All data is anonymised and meets with appropriate ethical standards for this type of community questionnaire.

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401 Figure Captions402

403 Figure 1. A) Responses to "Do you have a protocol for independent review of
404 quality-related results?"; B) Responses to "Do you have a written protocol for QA
405 review criteria?"

407 Figure 2. Average response to "Do you validate your project sample
408 measurements with: (Check all that apply)?"
409

410 Figure 3. Average responses to "What types of QC materials do you routinely use411 in analytical measurements for metabolomics projects? (Check all that apply)?











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