Solving CASMI 2013 with MetFrag, MetFusion and MOLGEN-MS/MS

Emma L. Schymanski,*1 Michael Gerlich,2 Christoph Ruttkies,2 and Steffen Neumann*2

1Eawag, Swiss Federal Institute of Aquatic Science and Technology, Überlandstrasse 133, 8600 Dübendorf, Switzerland
2Leibniz Institute of Plant Biochemistry, Department of Stress and Developmental Biology, Weinberg 3, 06120 Halle (Saale), Germany

The second Critical Assessment of Small Molecule Identification (CASMI) contest took place in 2013. A joint team from the Swiss Federal Institute of Aquatic Science and Technology (Eawag) and Leibniz Institute of Plant Biochemistry (IPB) participated in CASMI 2013 with an automatic workflow-style entry. MOLGEN-MS/MS was used for Category 1, molecular formula calculation, restricted by the information given for each challenge. MetFrag and MetFusion were used for Category 2, structure identification, retrieving candidates from the compound databases KEGG, PubChem and ChemSpider and joining these lists pre-submission. The results from Category 1 were used to guide whether formula or exact mass searches were performed for Category 2. The Category 2 results were impressive considering the database size and automated regime used, although these could not compete with the manual approach of the contest winner. The Category 1 results were affected by large m/z and ppm values in the challenge data, where strategies beyond pure enumeration from other participants were more successful. However, the combination used for the CASMI 2013 entries was extremely useful for developing decision-making criteria for automatic, high throughput general unknown (non-target) identification and for future contests.

Please cite this article as: E. L. Schymanski, et al., Solving CASMI 2013 with MetFrag, MetFusion and MOLGEN-MS/MS, Mass Spectrom (Tokyo) 2014; 3(3): S0036; DOI: 10.5702/massspectrometry.S0036

Keywords: computational mass spectrometry, small molecule identification, chemical clustering, maximum common substructure

INTRODUCTION

The Critical Assessment of Small Molecule Identification (CASMI) contest was inaugurated in 2012 to enable the comparison of structure elucidation methods using a common dataset and to enhance the exchange of identification approaches between disciplines.1) The liquid chromatography (LC-) tandem mass spectrometry (MS/MS) categories attracted the most participants, covering a wide variety of methods. In 2012, a combination of automated methods with expert knowledge was the winning strategy for Category 1 (molecular formula determination)2), although the de novo molecular formula predictions of MOLGEN-MS/MS3) and SIRIUS® also performed well.9) While the winner of Category 2 (structure identification) used database searching, the runners-up that used the relatively small compound database KEGG exhibited good ranks where the compound was present, at the cost of many cases where the correct answer was missing. Participants using much larger compound databases almost always had the correct solution present in submissions, albeit sometimes at very high ranks.10) In selected cases the molecules were small enough and with sufficiently detailed fragmentation patterns to allow structure generation approaches to be competitive: this even outperformed compound database strategies in three cases.11)

The CASMI 2013 contest led by Prof. Takaaki Nishioka included generally high resolution tandem mass spectra and fragmentation information for 16 challenge compounds, together with some biochemical background information.5,6) Figure 1 contains the structure and name of each challenge. The spectra were acquired on a variety of instruments and matrices, with varying accuracy and additional information (e.g. ion species identity and even formula in some cases—more details below), such that each challenge required some manual preprocessing and formatting. As for CASMI 2012,1) participants could submit molecular formulas (Category 1) or structural formulas (Category 2) for any or all challenges. Additionally, the organisers requested that participants indicate whether they used manual or automatic methods in preparing their submissions. Following the evaluation of the participant submissions and release of the challenge solutions, the organisers also declared some “highly commended” structures that they considered as being likely to produce spectral data very similar to the actual challenge data.5,6)

In light of the 2012 CASMI results and the information provided for the 2013 contest, preliminary decisions were made on the submission strategy. MOLGEN-MS/MS performed well for Category 1 (molecular formula prediction) in CASMI 2012 with a simple enumeration approach.5) Molecular formulas are generated combinatorially ac-

*Correspondence to: Emma L. Schymanski, Eawag: Swiss Federal Institute of Aquatic Science and Technology, Überlandstrasse 133, 8600 Dübendorf, Switzerland, e-mail: emma.schymanski@eawag.ch
*Correspondence to: Steffen Neumann, Leibniz Institute of Plant Biochemistry, Department of Stress and Developmental Biology, Weinberg 3, 06120 Halle (Saale), Germany, e-mail: sneumann@ipb-halle.de
According to the user inputs (elements, accuracy range and optional filters) and ranked according to the match to the MS1 isotope pattern, the MS2 fragment information (number of peaks with a sub-formula of the proposed formula) or a combined match value, as described in Meringer et al.\(^7\) Thus, MOLGEN-MS/MS was used for Category 1 entries for CASMI 2013, with the aim of incorporating this information in the Category 2 entries. The size of the challenges (up to 1,440 Da), relatively large ppm errors in some challenges and lack of diagnostic fragment information indicated that structure generation approaches would not be feasible for Category 2. Instead, the challenges favoured the compound database approaches of MetFrag\(^8\) and MetFusion\(^9\) for Category 2, molecular structure identification. MetFrag accepts a formula or mass plus accuracy information as an input to query compound databases such as PubChem,\(^{10}\) ChemSpider\(^{11}\) and KEGG\(^{12}\) for candidate structures. These are then fragmented \textit{in silico} using a combinatorial bond disconnection approach and scored according to the number of matching fragments in the fragment spectrum provided by the user, weighted by mass, intensity and the bond energy and ranked relative to the other candidates (see Wolf et al.\(^8\) for more details). In MetFusion, the results of the MetFrag ranking are combined with a mass spectral library search of the fragment spectrum, using the Tanimoto similarity to re-rank the candidates according to the spectral search information. For an unknown spectrum with similar spectra in the library, the scores are increased for similar candidates in the MetFrag results, while the results remain similar to those of MetFrag if no matching database spectra

**Fig. 1.** The challenge structures for CASMI 2013.
are present. More details are given in Gerlich and Neu-
mann.\(^{10}\) For CASMI 2013, the MOLGEN-MS/MS, MetFrag
and MetFusion approaches were combined with the aim of
markedly improving on the 2012 results\(^{3}\) and moving to-
towards a workflow-style automated elucidation.

**METHODS**

The development of a workflow approach for CASMI
2013 entries involved pre-processing the challenge data files,
creating query files for batch processing with MetFrag and
MetFusion with multiple databases for a broader chemical
coverage, then filtering and joining the results to automati-
cally create the submission files. Each run of the workflow
aimed to produce valid submission files and additional
constraints were added progressively following manual
inspection of the intermediate results. Several checks were
also added to ensure that the individual steps of the process
did not return wrong or incomplete results, in an “agile pro-
gramming” approach.

**Challenge data and pre-processing**

The challenge data for 2013 came from a variety of instru-
ments with different accuracies, isolation windows and vari-
ous measurement phenomena in the MS1 and MS/MS files.
The instruments (written as [challenge numbers] instru-
ment, ionisation, polarity abbreviation) were: [1–3] LCMS-
IT-TOF, ESI+; [4,5,9,10] LTQ Orbitrap Velos, nano-ESI+;
[6] APEX-II FT-ICR, ESI-; [7,8] Xevo QTOF, ESI+; [11,12]
LC-IT-TOF, ESI-; [13] AXIMA QIT MALDI+; [14] LCMS-
IT-TOF, APCI+; [15] Exactive Orbitrap and Xevo TQ, ESI+;
[16] Exactive Orbitrap and Xevo TQ, ESI-. Additional infor-
mation was also provided for some challenges and, where
possible, this was included in the automatic methods as
indicated below. More details on the measurements and the
additional information are available elsewhere.\(^{3,6}\)

A degree of pre-processing was necessary to apply a con-
sistent workflow on the files, while different parameters were
necessary for the different challenges to take full advantage
of highly accurate data when it was available. As neither
MOLGEN-MS/MS nor MetFrag/MetFusion account for iso-
tope patterns in the MS/MS, unequivocally identifiable iso-
tope peaks were removed from the MS/MS files (Challenges
1 & 2, 4–6, 9, 11, 14–16) to avoid incorrect formula/frag-
ment assignment to isotope peaks, especially problematic for MS/
MS where e.g. assignment to isotope peaks, especially problematic for MS/
1 & 2, 4–6, 9, 11, 14–16) to avoid incorrect formula/fragment
attempts were removed from the MS/MS files (Challenges
7 & 8 and 13 & 14. The mass accuracy settings for MS1 (“ppm”) and
MS/MS (“acc”) formula fitting, both in ppm, were (written as
[challenge number(s)] ppm&acc: [1–3] 10&10; [4] 5&10;
[5] 3&10; [6] 3&5; [9] 5&25; [10,15,16] 5&15; [11,12] 10&30.
The results were sorted by the combined match value, except
Challenge 15 which was sorted by MS/MS match value as no
isotope peaks were present in the MS1.

Generally, the challenges were calculated with the ele-
ments C, H, N, O, P and S (hereafter abbreviated to
CHNOPS), although additional restrictions were added ac-
cording to information provided for certain challenges\(^{9}\) or
as a result of manual inspection of the MS1 data and inter-
mediate results files. Challenge 1 and 2 formulas were fil-
tered according to RDB≤4 (additional information “aromatic
structures present”\(^{\text{a}}\)), while Challenge 3 formulas were
filtered according to C, N and O≥1 and RDB≥1 (additional
information “amide bonds present”\(^{\text{b}}\)). Elements for formula
calculations were restricted for Challenges 9 (5–15 C, 3 CI
10 (30–50 C) and 16 (5–20 C, 9–100 H, O≥2) using informa-
tion from the MS1 and MS/MS. For Challenge 15, the ppm
error in the MS/MS was used to separate formula scores,
with MS/MS peak scores scaled linearly from 1 to 0 between
5 and 15 ppm (all better than 5 ppm with score 1, all outside
15 ppm rejected completely). A detailed explanation of the
parameters in MOLGEN-MS/MS is available elsewhere.\(^{2,4}\)
For the submissions, the Category 1 results were cross-
checked with MetFrag results calculated with an exact mass
search and no element restrictions to see if any high-scoring
candidates with additional formulas were present.

For this manuscript, the molecular formula of all struc-
tural candidates were extracted from the MetFrag candidate
lists (via the InChI), together with the highest score per
candidate with the given formula. The formulas were then
ranked according to the score. These calculations were per-
formed for MetFrag using exact mass searches without any
element restrictions, MetFrag exact mass searches restricted
to compounds containing CHNOPS and also MetFusion ex-
act mass searches restricted to elements CHNOPS except for
Challenges 9, 15, and 16 (no restriction). These results were
then summarised across the databases to be comparable with
the joined datasets from Category 2. For the purpose of
concise display in this manuscript, the exact mass search
without element restrictions will be hereafter represented
as CHNOPS=F (where F stands for false) and with element
restrictions as CHNOPS=T (where T stands for true), con-
sistent with the MetFrag and MetFusion settings.

**Category 2: Structural formula**

MetFrag and MetFusion were used to retrieve candidate
structures from the compound databases PubChem,\(^{10}\)
ChemSpider\(^{12}\) and KEGG\(^{22}\) for submission as Category 2
tentries. The MetFrag parameters consisted of the exact mass
of the precursor ion (to search the databases within the er-
ger given, “ppm”) and the peak list retrieved from the chal-
gen data (for in silico fragmentation, within the given ppm
(parameter “mxppm”) and absolute boundaries (parameter

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\(^{1}\) Vol. 3 (2014), S0036
“mzabs,” in Da)). The parameters per challenge (written as [challenge numbers] ppm&nzmzppm&mzabs) were as follows: [1–3,7,8,11–14] 10&108&0.005; [4,5,9,10,15,16] 5&5&0.001; [6] 3&3&0.001. The in silico fragmentation was performed up to a tree depth of two. The ionization mode and charge were set for [M+H]⁺ or [M–H]⁻ as listed for Category 1. Further details about MetFrag and the scoring are given in Wolf et al.⁹ For challenges where the molecular formula was provided (Challenges 7, 8, 13 and 14) and where one formula was clear from the Category 1 results (Challenges 3, 4 and 9), the MetFrag candidate retrieval was performed using the formula instead of the exact mass.

MetFusion combines the in silico fragmentation approach of MetFrag with spectral database searching.⁹ The parameters for the MetFrag part of MetFusion were as described above. The spectrum search in MassBank required additional parameters. The list of instruments consisted of all ESI and additionally selected APCI and APPI instruments currently available in MassBank,¹⁵ including 21 different combinations of QqQ, QTOF, IT and ITIFT instruments. The ionization mode used in the spectrum search was chosen according to the information given with the challenges. The relative intensity cut-off was set to the MassBank default of 5, indicating that peaks with a relative intensity below that value are not examined in the peak matching algorithm of the spectrum search. MetFusion was set to limit the MassBank result list to 200 entries, which in general does not eliminate reference spectra that would improve the results. The database search was performed on the Japanese MassBank server. Both MetFrag and MetFusion kept all results from the compound database for a given query, only removing charged or disconnected compounds (e.g. salts). The candidates of the result lists obtained were filtered based on the first part of their InChI Key, to remove multiple stereoisomers of a given molecule (treated as equal in terms of MS).⁹

Additional submissions with different strategies were created for this manuscript. To assess the effect of database joining, candidate lists obtained from individual databases were evaluated separately, while all challenges were re-run with and without element restrictions to assess the formula filtering applied and, especially for Challenges 15 and 16, to generate results retrospectively that contained the correct candidate. Otherwise the settings and scoring were identical to the contest submissions.

Post-processing
After generating MetFrag and MetFusion batch query files for the individual databases, the results were joined to create the final submissions. All manipulations and plots were performed with scripts in the statistical language R,⁶ using the CDK,⁷ rCDK⁸ and OpenBabel⁹ for cheminformatics functionality. The database results were joined using the first block of the InChI Key, taking the highest scoring candidate from all databases. Charged structures that arose during post-retrieval interconversions were detected and removed (all structures containing \[7p or q]+ or \[-0–9\] entries, e.g. \[7p+1\] at the end of the InChI string). For challenges where more than one possible formula was selected specifically including the concept of “worst case rank” to evaluate candidates with equal scores. The worst case rank is the total of all candidates with better and equal scores compared with the correct candidate, so if an entry contains 5 candidates with higher scores and 2 additional candidates with the same score as the correct candidate, the best case rank would be 6 and the worst case rank 8.

To facilitate the assessment of the scored candidate lists, a hierarchical clustering based on the chemical similarities between candidates was performed. This representation allows a quick assessment of structural similarity, as well as indicating whether clusters contain only individual candidates, or several consistently high-scoring candidates. In addition, the maximum common substructure (MCSS) of all candidates contained in a cluster provided information about the scaffold of the candidates. Additional visualisations included 1) the structural “distance” between the correct solution and the top ranking candidates, represented by the Tanimoto similarity,¹¹ 2) the overlap of the MCSS from each cluster and the correct solution using AMBIT²⁰ and 3) the peaks of the MS/MS spectrum expected by MetFrag overlaying the experimental spectrum. The resulting plots are given in the Supporting Information (SI), merged and numbered sequentially per challenge, as well as for selected challenges below. Explanations to assist interpreting the plots can be found on the first page of the SI.

**RESULTS AND DISCUSSION**

**Category 1**

The calculations for MOLGEN-MS/MS were generally very quick and were under 1 s for all challenges except Challenges 15 (2 s), 6 (94 s) and 10 (1,027 s) on an Intel® i5 CPU M560 with 4 GB memory. However, the results for Category 1 reflected the difficulty of using a purely combinatorial approach in molecular formula determination for compounds with large masses. These are summarised in Table 1 (column “Cat. 1 Entries”), along with the precursor mz and the formula. The results with MOLGEN-MS/MS were generally good (rank 1 or 2) for compounds with small mz precursors, but no longer competitive for larger mz values where other CASMI participants with different approaches outperformed MOLGEN-MS/MS.⁵,⁶ The winner of Category 1 ranked the correct formula in first place for all challenges.

Pre-submission, the MOLGEN-MS/MS results were cross-checked with preliminary Category 2 results using MetFrag with an exact mass search to see if any high-scoring candidates had different formulas to those proposed by MOLGEN-MS/MS, which could indicate that the element restrictions used were incorrect. These results are shown in the fifth column of Table 1, sorted using the highest-scoring candidate in the MetFrag results list with the given formula. The correct formulas for Challenges 15 and 16 were ranked 22nd and 4th, respectively, and as a result, F-containing formulas were not considered for these challenges (incorrectly in hindsight). In 9 of 12 challenges, taking the highest-scoring MetFrag candidate per formula to rank the formula resulted in an equal or better rank than that achieved with MOLGEN-MS/MS. Challenge 5 was an exception, however
restricting the elements to C, H, N, O, P and S only (6th column of Table 1) resulted in likewise 9 of 12 challenges with equal or better results than MOLGEN-MS/MS, but losing the correct formulas for Challenges 9, 15 and 16, which all contained halogens. The retrospective results were more remarkable with MetFusion. If the Category 1 submissions had been made using the formula from the top scoring MetFusion candidates, all challenges would have had the correct formula ranked in first place, including no element restrictions for Challenges 9, 15 and 16. This is shown in the last column of Table 1.

In retrospect, a workflow approach for unknown identification on the basis of database queries may be performed efficiently using an exact mass search with post query formula filtering to ensure that all possible candidates are captured and considered. However, the drawback of such an approach is that database queries, even of large databases, will never ensure 100% coverage of chemical space and thus the results of formula generation should not be discounted completely. To demonstrate this, if only KEGG was used for the database queries, 8 of the 12 formulæ would have been correctly ranked 1st, but the remaining 4 challenges would have had no candidate formulæ. Thus, true non-target identification will always require formula calculation to ensure that the results are not only restricted to database entries.

### Category 2

The results submitted for CASMI 2013 in Category 2 are shown in Table 2. The calculation times for MetFrag and MetFusion per challenge, including database retrieval, ranged from less than one minute for challenges with few candidate structures up to 30 min for challenges with several thousand candidates. The runtime depends on many factors such as the speed of the online query, settings, number of candidates and fragments to match as well as the system configuration. The post-processing took just under 6 hrs for all challenges at this conceptual stage, with interconversions between representations the most time-consuming. Calcula-

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### Table 1. Summary of Category 1 data and results, including the CASMI 2013 MOLGEN-MS/MS submissions and retrospective entries compiled using MetFrag and MetFusion with exact mass searches with (CHNOPS=+T) or without (CHNOPS=-F) element restrictions. Results are Rank (Total Candidates).

| Chal. | Precursor m/z* | Formula | MOLGEN-MS/MS Entries | MetFrag CHNOPS=F | MetFrag CHNOPS=T | MetFusion CHNOPS=TI/F |
|-------|----------------|---------|----------------------|----------------|----------------|---------------------|
| 1 (+) | 314.1364       | C₆H₄NO₄ | 2 (11)               | 1 (44)         | 1 (13)         | 1 (13)              |
| 2 (+) | 265.1524       | C₅H₄N₂O₄ | 1 (2)                | 1 (13)         | 1 (7)          | 1 (7)               |
| 3 (+) | 301.1843       | C₅H₅N₂O₄ | 1 (3)                | 1 (21)         | 1 (9)          | 1 (9)               |
| 4 (+) | 211.1121       | C₃H₇O     | 1 (4)                | 1 (6)          | 1 (4)          | 1 (4)               |
| 5 (+) | 291.0719       | C₄H₅O₂S   | 1 (9)                | 4 (46)         | 1 (10)         | 1 (10)              |
| 6 (-) | 935.6224       | C₈H₆O₆P   | 8 (144)              | 1 (9)          | 1 (9)          | 1 (7)               |
| 9 (+) | 349.9337       | C₆H₆ClNO₃PS | 1 (6)              | 1 (16)         | –              | – /0 (1) /1 (15) |
| 10 (+) | 922.5351      | C₆H₇N₃O₉₁ | 45 (283)             | 3 (10)         | 3 (7)          | 1 (3)               |
| 11 (+) | 337.1080      | C₃H₃O₆   | 4 (41)               | 1 (55)         | 1 (21)         | 1 (21)              |
| 12 (-) | 269.0449      | C₃H₇O₆   | 9 (27)               | 2 (40)         | 2 (12)         | 1 (12)              |
| 15 (-) | 462.9974      | C₅H₅F₄O   | – (177)              | 22 (105)       | –              | – /7 (1) /6 (9) |
| 16 (+) | 362.1527      | C₆H₇F₃NO₄ | – (6)                | 4 (51)         | –              | – /15 (1) /5 (6) |

*Exact mass of ion (i.e., [M+H]+ or [M–H]−).*

### Table 2. Category 2 results as submitted using database joining, compared with MetFrag & MetFusion with individual databases. Columns include “Worst Case Rank (Total Candidates).”

| Challenge | CASMI Entry | MetFrag ChemSp* | MetFrag KEGG² | MetFrag PubChem* | MetFusion ChemSp* | MetFusion KEGG² | MetFusion PubChem² |
|-----------|-------------|-----------------|---------------|----------------|-----------------|----------------|-------------------|
| 1         | 9 (5633)²C  | 1124 (4247)     | 1 (4)         | 1078 (5479)    | 7 (4064)        | 1 (4)          | 9 (4723)          |
| 2         | 4 (12702)²C | 107 (3244)      | 1 (1)         | 649 (11925)    | 20 (3228)       | 1 (1)          | 34 (11566)        |
| 3⁺        | 21 (335)²B  | 1 (52)          | – (–)         | 21 (325)       | 2 (52)          | – (–)          | 15 (325)          |
| 4⁺        | 238 (721)²B | 6 (323)         | – (4)         | 7 (705)        | 90 (323)        | – (4)          | 233 (705)         |
| 5         | 4 (366)²C   | 9 (891)         | 1 (1)         | 39 (2303)      | 3 (258)         | 1 (1)          | 4 (593)           |
| 6         | 1 (6)       | – (5)           | – (–)         | 1 (1)          | – (5)           | – (–)          | 1 (1)             |
| 7⁺        | 17 (17)²B   | 16 (16)         | 2 (2)         | 17 (17)        | – (–)⁹          | – (–)³         | – (–)³           |
| 8⁺        | 1 (1)       | 1 (1)           | – (–)         | 1 (1)          | 1 (1)           | – (–)          | 1 (1)             |
| 9⁺        | 1 (4)²*     | 1 (2)           | 1 (1)         | 2 (4)          | 1 (2)           | 1 (1)          | 1 (4)             |
| 10        | 1 (9)       | 4 (14)          | – (–)         | 6 (24)         | 1 (3)           | – (–)          | 1 (9)             |
| 11⁺       | 1 (2392)    | 2 (2317)        | 1 (10)        | 15 (3843)      | 1 (1430)        | 1 (9)          | 1 (2204)          |
| 12        | 35 (902)    | 149 (561)       | 15 (18)       | 207 (1103)     | 38 (438)        | 9 (18)         | 41 (813)          |
| 13⁺       | 12 (227)    | 11 (78)         | – (–)         | 17 (210)       | 7 (78)          | – (–)          | 13 (210)          |
| 14⁺       | 1 (8219)    | 151 (4793)      | 4 (10)        | 250 (7307)     | 71 (4793)       | 1 (10)         | 146 (7308)        |
| 15⁺       | – (–)⁶C     | – (–)           | – (–)         | – (16)         | – (–)           | – (–)          | – (–)             |
| 16⁺       | – (–)⁶     | 135 (449)       | – (–)         | 179 (663)      | 1 (127)         | – (–)          | 1 (401)           |
| 16⁻       | – (–)⁶     | – (–)           | – (–)         | – (6955)       | – (4270)        | – (–)          | – (–6633)         |
| 16⁻       | – (–)⁶     | 5251 (5251)     | 1 (1)         | – (8855)       | 1 (3101)        | 1 (1)          | 1 (5071)          |

*Best tautomer results reported. ²Restricted by formula. ³CASMI entry only restricted by Category 1 results (see text for details). ⁴Results with CHNOPS=True. ⁵Results with CHNOPS=False. Note for Challenges 15 and 16 the letter in the row name takes precedence over the column header. ⁶No peaks found in MassBank. ⁷CASMI submission performed using MetFrag, not MetFusion. Bold type in "CASMI Entry" indicates challenge was equal or outright winner.
tions were performed on a computing server, equipped with two 6176 AMD Opteron™ Processors (24 cores) and 96 GB RAM for efficient parallel computations and multi-user configuration.

In total these submissions were the (joint) winner in 7 of the 16 challenges, where 6 of these had the correct compound ranked number 1—including two challenges (11, 14) with over 2,000 possible candidates. The joining of the three database results meant that the correct entry was present in all submissions despite the fact that 7, 1 and 1 of the correct entries were missing in KEGG, ChemSpider and PubChem, respectively. Thus, the automatic database joining avoided the coverage problem experienced by participants who relied on PubChem alone, and approached the manual strategy to include many data sources applied by the winners. 5,6)

In addition to the contest submissions, the results that would have been achieved with either MetFrag or MetFusion alone on the individual databases are shown in the remaining columns of Table 2, as well as additional rows for Challenges 15 & 16 with an exact mass search and no element restrictions, to demonstrate what could have been achieved for these challenges. As expected, MetFusion outperformed MetFrag, with better or similar ranks for all challenges except Challenges 4 and 7. For Challenge 4, many candidates with high MetFrag scores were also structurally similar to the best spectral match in MassBank (Resveratrol, accession number CE000096, spectral similarity 0.5), which elevated the scores for these candidates above the score of the correct structure. No peaks corresponding to the spectrum of Challenge 7 were found in MassBank, such that only MetFrag results were available. As seen in Table 2, Challenges 15 and 16 were a missed opportunity, as running MetFusion with an exact mass search and no element restrictions would have resulted in two additional wins with the correct compound ranked first. Further details are given per challenge below.

**Challenge 1:** The Category 2 entries for Challenge 1 were restricted to the formulas C_{14}H_{15}N_{7}O_{2} and C_{18}H_{19}NO_{4} using information from Category 1. MetFusion improved the results dramatically over MetFrag, where the results contained many candidates with the same scores. The maximum common substructure (MCSS) of the cluster containing the correct candidate (cluster 13, see the Supporting Information (SI), Figure S1) covers the majority of the backbone, excluding only one methoxy and hydroxy substituent per aromatic ring.

**Challenge 2:** This challenge had the most candidates of all challenges and although formula pre-selection was performed (candidates with C_{14}H_{15}N_{7}O_{2} or C_{10}H_{16}N_{8}O), the vast majority of candidates had the former (correct) formula. Challenge 2 was one of the only two challenges where the joining of the databases resulted in a worse rank than the worst of the individual database results (see Table 2; the other was Challenge 4). Both challenges had reasonably

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**Fig. 2.** Challenge 2. (a) Cluster plot and (b–e) selected maximum common substructures for Challenge 2. Cluster 17 is the correct structure, feruloyl putrescine.
large candidate numbers and very little difference between the scores (Challenge 2 scores 0.7350–1.0885, with differences between the candidates in the 2nd decimal place). These are cases where the measured spectra and resulting spectral database matches do not provide enough discrimination power between all candidates. Although MetFrag explains the three most intense peaks in both Challenge 2 (Figure S2, SI) and 4 (Figure S4, SI), this is not sufficient to separate such a large number of candidate structures. Despite this, the rank of 44 of 12,702 candidates is a good result for automatic methods. A cluster plot of the top 50 compounds (Fig. 2a) shows that the correct structure forms a single-compound cluster (Fig. 2d), which is very different to the maximum common substructure from the cluster of the top scoring compounds (Fig. 2c) and other clusters containing high-scoring candidates (Figs. 2b & c). Thus, for this challenge the clustering would not even help to isolate a compound class, supporting the conclusions above that the measured spectrum does not provide sufficient information to rank so many candidates.

**Challenge 3:** These entries were restricted to candidates with the formula C_{13}H_{24}N_{4}O_{4} using the Category 1 results. The MetFrag and MetFusion results were very similar for this challenge, indicating a lack of database spectra in MassBank for these compounds (highest spectral similarity 0.26). Submissions were thus made using MetFrag, which explained the two most intense peaks (see SI, Figure S3), the third was very low intensity. The CASMI 2013 organisers explained the two most intense peaks (see SI, Figure S3), which resulted in a worst case rank of 21 for both, while a best case rank would equal to 1 for both structures. The clustering and similarity plot shows the relationship between the two structures impressively, with the two clustering together and similarity plot shows the relationship between the two structures impressively, with the two clustering together and clearly separated in similarity from the rest of the candidates (Figure S3, SI). Again, as few fragments were present in the measured spectrum, the high-scoring candidates were distributed amongst many clusters.

**Challenge 4:** was a disappointing result where MetFusion made the result worse than MetFrag, which is surprising for a reasonably simple structure where MetFrag covered the predicted spectrum reasonably well (see SI, Figure S4). Only candidates with the formula C_{13}H_{14}O_{4} were considered. This challenge also had highly commended alternative solutions, which were both ranked lower than the correct solution (ranks 293 and 299 for 2-(2-methylphenyl)-1-phenylethanolone and 2-(4-methylphenyl)-1-phenylethanolone, respectively). Interestingly, these two highly commended structures group together with some other similar compounds in the similarity plot (SI, Figure S4).

**Challenge 5:** was another challenge with a highly commended structure; the original submission had the correct structure isoprothiolane ranked 4th (Fig. 3b) and the highly commended structure, dipropyl 1,3-dithiolan-2-ylidenemalonate, was ranked 1st (Fig. 3c). These two structures cluster together in Cluster 30 with a very good MCSS (Fig. 3a) and are also clearly separated from all other candidates in the similarity plot (SI, Figure S5). Many single-compound clusters are present for this challenge, showing the diversity of candidates. Formula preselection was performed using all 9 formulas present in the Category 1 submission.

**Challenge 6:** was a very successful challenge with the correct structure ranked first of six candidates, without applying any formula preselection. Each candidate was member of a single-compound cluster, as shown in the SI, Figure S6.

**Challenge 7:** The formula was provided for this challenge (C_{12}H_{20}O_{10}) and as there were no matching peaks in MassBank, the submission was performed using MetFrag. Although the overall result looks poor (17th of 17), this is only a result of the CASMI evaluation using the “worst case rank” and in fact all candidates had equal scores. The correct solution is a flavonoid pentamer, the challenge spectrum shows major peak groups every 288Da (C_{15}H_{12}O_{6}), with MetFrag explaining one peak in every major peak group. Minor peak groups represented a loss of 152Da (C_{15}H_{12}O_{6}). Challenge 7 is a very large molecule and all 17 candidates are very similar, manual interpretation indicates that all candidates explain the challenge spectrum equally well, as shown in Figure S7 (SI).

**Challenge 8:** This challenge was a trivial case, with one (correct) candidate in both PubChem and ChemSpider with the given formula C_{6}H_{10}O_{3}PSCl to result in a rank 1 of 1.

**Challenge 9:** The candidates for this challenge were restricted to C_{6}H_{10}NO_{3}PSCl and submissions were made using MetFrag. Although only a minor part of the spectrum was explained (see Figure S9, SI), the O,O,O-substituted phosphorothioates (score>0.6) were clearly separated from the O,O,S-substituted phosphorothioates (score <0.2). Although the top two isomers actually had the same MetFrag score, one candidate (chlorpyrifos) is much more common than the other candidate (supported by the number of references in ChemSpider, 1,457 versus 4) and a spectrum was available internally (also now on MassBank, accession numbers EA295001–EA295014, with a spectral match of 0.92). On the basis of this evidence, 0.5 was added to the score of chlorpyrifos.

![Fig. 3. Challenge 5.](image)

(a) MCSS for Cluster 30. (b) The correct structure, isoprothiolane, overlaid with the MCSS from Cluster 30 (which covers either terminal methyl group). (c) The highly commended structure, dipropyl 1,3-dithiolan-2-ylidenemalonate overlaid with the MCSS from Cluster 30.

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rifos manually, resulting in a final rank of 1. The MCSS and two top structures are shown in Fig. 4, while the cluster plot is given in the SI.

**Challenge 10:** This was another good result, where most participants in CASMI 2013 picked the correct structure at rank 1. MetFusion improved the good MetFrag result further, as the majority of the peaks were explained by MetFrag. No formula restriction was used for this challenge.

**Challenge 11:** No formula restrictions were used for this challenge, which had three equally valid solutions in the end due to the presence of different tautomeric forms of challenge 11 in the compound databases.5,6) The different location of the single and double bonds meant that in silico methods and databases regarded these structures differently and the three participants who submitted entries for these challenges reported either two or three of the tautomers, each with different ranks.6) MetFrag clarified two peaks more of the spectrum with the first tautomer (Figure S11, SI), which was ranked first and thus a winning entry (shown in Fig. 5a). The other tautomers were ranked 21st and 22nd (Figs. 5b and c) and cluster together to form a much bigger MCSS than the cluster containing the winning tautomer (Figure S11, SI). Interestingly, the change in resonance form has a large impact on the similarity calculation, the tautomer ranked in first place only has a similarity of ~0.65 to the other two tautomers (see Figure S11). Despite this, the MCSS for the major clusters show that most candidates have similar connectivity and if the MCSS were less sensitive to resonance structures, all tautomers could be encompassed into a general MCSS with the SMARTS c1cc(O)ccc1C= C—C(=O)—C—C(=O)—C=Cc2ccc(O)c2.

**Challenge 12:** Challenge 12 was a reasonably solid result, with rank 35 of 902 candidates, without any formula restrictions to the candidate lists. MetFrag clarified most of the high intensity peaks (see SI Figure S12) and the similarity plot shows that the top 40 compounds are all very similar (>0.75). With many similar structures amongst the candidates, this rank is a reasonable result and a more detailed manual analysis, such as that performed by other participants, would be required to find the correct structure.

**Challenge 13:** This was the one challenge where the MetFrag/MetFusion submission was the only winner, with a rank of 12 (of 227) compared with 24 and 42 for the other two teams participating in this challenge, using the formula C_{17}H_{30}N_{2}O_{5} provided. In this case both MetFrag and MetFusion returned very similar results as there were no very similar spectra in MassBank (highest match 0.58 for Etodolac, CO000187). MetFrag only clarified one of the 7 peaks, while both the cluster and similarity plots show that the candidates for this challenge were very diverse, with many single-compound clusters (see Figure S13, SI). As such, it would be difficult to define a substance class using automated methods for this challenge, similar to Challenge 2.

**Challenge 14:** This challenge had an exceptional result of top rank of 8219 candidates with the formula C_{21}H_{24}N_{2}O_{3} (given), where the joining of database results had a big influence on the outcome. Like Challenge 2, there is little differentiation in the scores, visible in the ChemSpider and PubChem ranks of 71 and 146, respectively. However, the top two KEGG candidates had higher scores (0.7148 and 0.7132) than all PubChem and ChemSpider candidates (top scores 0.7112 and 0.7091, respectively). The cluster plots show that
the top six compounds cluster into two groups, again with a large MCSS that explain a large proportion of the molecule. A very similar compound with rank 8 is also included in the cluster with the correct molecule, these three compounds are clearly separated in the similarity plot (Figure S14, SI), while the compounds in the second cluster form the next group of similar compounds (ranks 3–6) in the similarity plot. This excellent result for MetFusion is due to a high matching MassBank spectrum for Ajmalicine (FJO00003), a stereoisomer of the correct answer with a spectral similarity score of 0.8.

**Challenge 15:** As mentioned above, Challenges 15 and 16 were missed opportunities as MetFrag was used to see if high scoring candidates with different formulas to the pre-selected formulas were present. MetFusion results were then submitted using the CHNOPS-T filter (no further formula restrictions). The poor performance of MetFrag for both challenges can be seen easily looking at the overlay of measured and predicted spectra (SI, Figure S15). However, the (retrospective) MetFusion results in Table 2 show that both these challenges would have been ranked number 1 if MetFusion had been used with exact mass and no element restrictions. For Challenge 15 the correct candidate had a MetFrag score of 0.26675, which was raised to 1.00739 after weighting with MassBank, due to a high structural similarity (0.78) between the correct answer and the best and high-scoring spectral match, 82Fluorotelomer Acid (FF000410).

**Challenge 16:** While the original CASMI submission did not contain the correct compound, retrospective calculations were performed using an unrestricted exact mass search to obtain results containing the correct compound. These calculations showed that the original MetFrag score of 0.489 for the correct candidate increased to 1.526 after MetFusion due to a spectral match of 0.69 with the correct compound Ofloxacin in MassBank (UF407501), which elevated the correct candidate into first place. This clearly shows the benefits of adding spectra of standard compounds into openly accessible spectral libraries and also the strength of the combined approach of MetFrag and MetFusion. The clustering results shown in Figure S16 (SI) were also interesting and despite the very high similarities with many top candidates, informative MCSS were formed to allow a good separation of candidates.

The results from the challenges above show that the approach used here was generally very successful for an automated approach using large compound databases and the results were a great improvement on the compound database submissions for the 2012 CASMI competition. The implementation of filtering criteria post-candidate retrieval proved an efficient and also flexible means of reviewing the results and preparing the desired submission files. Prior determination the molecular formula and then searching on that formula only is generally performed in non-target identification and can reduce the number of higher scoring incorrect candidates. However, restricting the submissions to one or a few formulas upfront can result in missing solutions as seen in Challenges 15 and 16. For those challenges where the formula was not clear, the exact mass search with post-retrieval filtering for selected formulas certainly proved a viable alternative. MetFusion was also often able to rank the correct candidate highly even without formula restrictions. As mentioned above, the joining of the databases improved coverage while lowering the ranks below the lowest rank for an individual database in only two cases. This may be partially due to the score normalisation implemented in individual MetFrag and MetFusion runs, which is an issue being investigated beyond the scope of CASMI 2013. Other features that would be useful in the future to improve the results further would be ways to incorporate more of the additional information provided by the organisers, such as “the compound is a secondary metabolite” (Challenge 1) or “this is a synthetic product” (Challenge 3). While some research has been done into indexes of metabolite likeness (e.g. Hamdalla et al., Petroncely et al.) or using references to rank candidates (Little et al.), these features were not yet implemented into MetFrag or MetFusion at the time of CASMI 2013.

While the workflow approach here was generally successful, a number of anomalies exist. The interconversions between various cheminformatic data formats (mol, SDF, SMILES, InChI) during the workflow can lead to changes in the structures and the location of hydrogens, especially where aromatic nitrogens are involved. The MCSS in several cases seemed to be much smaller than necessary due to inconsistencies in aromatic and/or tautomeric perception. Large compound databases inherently have errors, which any workflow will need to catch or work around (e.g. PubChem CID: 59694784 retrieved for Challenge 2, but with a CH group instead of CH₂, or PubChem CID 24209464 for Challenge 4 which does not generate valid fingerprints for clustering). As observed by the CASMI organisers for Challenge 11, tautomeric structures and other chemically equivalent forms of structures that are considered different informatically but interchangeable chemically, pose a problem for any workflow especially as the different representations can generate different ranks. The assumption made during MetFrag/MetFusion and in the post-generation filtering to use the InChI Key first block (i.e., not considering stereochemistry) holds reasonably well for mass spectral-based methods, where the stereochemistry cannot be determined (unless e.g. chiral chromatography or ion mobility is used, i.e., via separation, not detection). This sometimes (but not always) captures tautomers and the issue of tautomerism should be investigated closer in the future. While the post-filtering options applied were very efficient at removing some undesired phenomena in the results lists (e.g. charged structures), it is unlikely that this affected the ranks in the CASMI contest dramatically as these were generally low scoring candidates. Although the total candidate numbers were reduced, the absolute rank is what matters in the end for identification.

**CONCLUSION AND PERSPECTIVES**

The aim with the submissions for CASMI 2013 presented here were two-fold: to improve on CASMI 2012 results and make progress towards a high throughput identification workflow on the basis of MetFrag and MetFusion. Both aims were successful and although the Category 1 results were disappointing compared with the other participants, this did not hinder the results for Category 2, which were very good considering the large compound databases used. Although the strategy of joining three databases increased the candidate numbers, it also increased the coverage of the submissions and in only two cases lowered the rank below...
the lowest rank of the largest individual database. As none of the three databases used contained all 16 challenges, the joining of results from several databases is certainly advantageous. Combinatorial molecular formula prediction, even with MS1 and MS2 fragment scoring, clearly requires additional restrictions for large m/z values. In these cases, unrestricted elements and an exact mass retrieval combined with post-scoring filtering via formula selection proved a viable alternative to upfront formula determination. Although taking the top formula from the MetFusion ranks would have resulted in a number 1 ranking for all CASMI 2013 challenges, this is unlikely to hold for a greater number of cases and this “success rate” would be worth investigating on a larger dataset of known compounds. As it is possible that true unknown compounds are being considered in genuine non-targeted studies (rather than in the current contest-style study), consideration must be given to compounds that are not necessarily in a compound database. In these cases a molecular formula calculation must be considered and full elucidation would have to move beyond compound database queries alone. Further improvements to a high-throughput formula determination would be the greater inclusion of restrictions, such as inclusion of elements detected via isotope patterns (Cl, Br) or isotope fine structure for high resolution instruments (\(^{18}\text{O},^{13}\text{N}\)) and even substructure information such as the aromaticity and amide bond information given for Challenges 1–3.

Acknowledgements

We would like to thank Prof. Nishioka and the CASMI 2013 organisers, as well as all the other participants and all contributors to open software, databases and spectral libraries used in this work. Comments from Juliane Hollender and Stephan Schmidt improved the manuscript. CR and MG acknowledge funding from DFG Grant NE/1396/5-1, while ES acknowledges funding for the European Union for the Marie Curie Inter-European Postdoctoral Fellowship 299734.

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