Software-assisted, high-throughput identification of main metabolites of pharmaceutical drugs

Rapid data acquisition by Agilent 1290 Infinity LC, TOF and Q-TOF instrumentation, and subsequent identification of metabolites by Agilent MassHunter Metabolite Identification software

Application Note

Metabolite identification in drug discovery and drug development

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Abstract
This Application Note describes:
• Rapid separation of metabolites generated from in-vitro experiments using the Agilent 1290 Infinity LC, system
• Fast acquisition of TOF mass spectra using Agilent 6530 Accurate-Mass Quadrupole Time-of-Flight LC/MS systems
• Fast, software-assisted identification of main metabolites from in-vitro experiments using Agilent MassHunter Metabolite Identification software
• Generation of reports for the identified metabolites using Agilent MassHunter software
**Introduction**

In modern pharmaceutical drug development it is of crucial importance to analyze the adsorption, distribution, metabolism and excretion (ADME) properties of possible new drug candidates as quickly as possible in order to make decisions about further investments in the development of a special compound. To find compounds with the correct properties it is essential to screen a large number of compounds for their ADME properties, which requires to work in an high-throughput environment. This Application Note describes the application of the Agilent 1290 Infinity LC system, the Agilent 6530 Q-TOF MS system and the MassHunter Metabolite Identification software for fast, high-throughput identification of main metabolites of new pharmaceutical drug candidate compounds.

**Experimental**

**Equipment**
- Agilent 1290 Infinity LC system consisting of 1290 Infinity Binary Pump with integrated degasser, 1290 High Performance Autosampler with thermostat, and 1290 Infinity Thermostatted Column compartment
- Agilent 6530 Accurate-Mass Q-TOF LC/MS system
- Agilent MassHunter Metabolite Identification (MetID) software
- Column: ZORBAX SB-C18, 2.1 x 50 mm, 1.8 µm

**Sample preparation**
The following stock solutions were used:
- 20 mg/mL microsomal S9 preparation
- 0.1 mg/mL buspirone in water
- 1.6 mg NADP in 1.6 mL 0.1 M phosphate buffer, pH 7.4
- 50 mM isocitrate/MgCl$_2$ (203 mg MgCl$_2$·6H$_2$O + 258.1 mg isocitrate in 20 mL H$_2$O)
- Isocitrate dehydrogenase 0.33 unit/µL
- NADPH regeneration system: 1.6 mL NADP solution + 1.6 mL isocitrate solution + 100 µL IDH solution.

**LC method**
- Solvent A: Water + 0.1 % formic acid
- Solvent B: ACN + 0.1 % formic acid
- Flow: 0.8 mL/min
- Gradient: 0 min, 5 %B; 0.10 min, 5 %B; 1.10 min, 75 %B;
- Stop time: 1.10 min
- Post time: 1 min.
- Injection: Volume 5 µL, sample cooler at 4 °C, needle wash in 50 % methanol for 5 s, injection loop to bypass at 0.1 min with flush out factor 16
- Column: Temperature 60 °C

**TOF MS method**
- Source: ESI positive
- Capillary: 3500 V
- Dry gas: 12 L/min
- Nebulizer: 55 psi
- Gas temp.: 350 °C

**Data analysis method in the MetID software**
The first step in the analysis comprised a comparison between the data file that contained the metabolite compounds (metabolite sample) and the data file that contained only the parent drug (control sample). All detectable mass signals were extracted from the MS level data using the Molecular Feature Extraction (MFE) algorithm. Related compound isotope masses and adduct masses were grouped together into discrete molecular features, and chemical noise was removed. The compounds lists of the metabolized sample and the control were then compared.

All new compounds or those that increased twofold in the metabolized sample were considered potential metabolites and were subjected to further analysis by different algorithms. The algorithms can identify and qualify new metabolites, or just qualify metabolites found by another algorithm. In this high-throughput experiment all algorithms’ results were weighted equally and combined into a final identification relevance score. Metabolites were qualified when their final score was above the stringently defined relevance threshold. The results from all algorithms were collated in a results table, which could be inspected at-a-glance and reported.
Results and discussion

To achieve fast separation of the metabolites on a 50 mm, 1.8 µm particle size column, a 1 minute gradient was applied by the Agilent 1290 Infinity LC system. The metabolites were generated from the pharmaceutical test compound buspirone in an in-vitro assay. For adequate detection with the time-of-flight mass spectrometer the instrument was operated at a data rate of 5 Hz.

After generation the data was loaded into the MetID software and analyzed using a common method. The result was displayed by the MetID software in an at-a-glance table, in which the result for each metabolite could be examined in more detail (figure 1). From the results table a summary report was generated, which showed the available information for each metabolite (figure 2). The more extensive report contained the detailed results for each metabolite. As example the result for a mono-hydroxyl metabolite (figures 3 to 5) and a dihydroxy metabolite (figures 6 to 8) of buspirone are discussed here.

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<th>Mass</th>
<th>RT</th>
<th>Rel.</th>
<th>Qual.</th>
<th>User</th>
<th>SC</th>
<th>IPM</th>
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Figure 1
Result table showing an at-a-glance summary of buspirone metabolite analysis with overall identified metabolites, extracted ion chromatograms (EIC), extracted compound chromatograms (ECC), isotopic pattern analysis and calculated formulas.

Figure 2
Summary result report, including qualified metabolites sorted by their retention times (RT), with their metabolite names and relative score, molecular mass and the passed flag for individual algorithm results. SC=Sample-control comparison, IPM=Isotopic Pattern Matching, EIC=Extracted Ion Chromatogram, MDF=Mass Defect Filter, Form.=Calculated Formula, BioXF=Assigned Biotransformation, Qual.=Qualified by Score, User=Qualified by User.
The extensive report for the mono-hydroxyl metabolite, which eluted after 0.75 minutes at m/z 402.2511, showed the detailed information about the metabolite itself such as measured accurate mass, calculated formula, assigned biotransformation and ion species. Further, the report showed more detailed information about the result of each individual algorithm, for example, Molecular Feature Extraction (MFE), Extracted Ion Chromatogram (EIC) compound search and Mass Defect Filter Result (figure 3). For the hydroxyl metabolite the possible formula was calculated based not only on a defined mass error window but also on the measured isotopic pattern, which increased the quality of the calculated formula and limited the possible number of hits significantly. These results were also displayed in the detailed metabolite result report for the formula (figure 4).

Figure 3
Detailed metabolite report for the buspirone hydroxy metabolite at retention time 0.75 min. This part of the report gives detailed information about the identified metabolite and the identifying algorithms. Other detailed information about formula (figure 4), chromatograms and isotopic pattern (figure 5) are also available.

Figure 4
Detailed metabolite report about the formula including isotopic pattern, calculated for the buspirone hydroxy metabolite at retention time 0.75 min.
Finally, the EIC, ECC and isotopic pattern were displayed (figure 5). The EIC of m/z 402.25 showed 5 peaks for possible hydroxyl metabolites of buspirone with the selected one at retention time 0.75 minutes (figure 5A). The ECC showed the extracted MFE compound for the molecular mass of 401.2439 at retention time 0.75 minutes identical to the EIC (figure 5B). The measured isotopic pattern of this compound showed an excellent fit to the calculated isotopic pattern as a basis for the formula calculation (figure 5C).

Within the same data analysis the dihydroxy metabolites at a level of two orders of magnitude below the mono-hydroxy metabolites were also identified. The extensive report showed detailed information about the dihydroxy metabolite, which elutes after 0.71 minutes at m/z 418.2461 and the detailed information about each algorithm (figure 6).

**Figure 5**
Detailed metabolite report for buspirone hydroxy metabolite at retention time 0.75 min:
A) Extracted Ion Chromatograms (EIC) of compounds with mass 402.25
B) Extracted Compound Chromatogram (ECC) of buspirone hydroxy metabolite at retention time 0.75 min
C) Measured isotopic pattern of buspirone hydroxy metabolite at retention time 0.75 min (blue lines) and calculated isotopic pattern (CIP, green box).

**Figure 6**
Detailed metabolite report for dihydroxy metabolite of buspirone at retention time 0.71 min. This part of the report gives detailed information about the identified metabolite and the identifying algorithms. Other detailed information about formula (see figure 7), chromatograms and isotopic pattern (see figure 8) are also available.
The calculation of the formula was outlined in the detailed formula report (figure 7).

The EIC of m/z 418.24 showed about five significant peaks for possible dihydroxylated metabolites of buspirone with the selected peak at 0.71 minutes (figure 7A). The ECC showed the extracted MFE compound for the molecular mass of 417.2388 at retention time 0.71 identical to the EIC (figure 7B). The measured and calculated isotopic pattern of this compound is shown in figure 7C.

**Metabolite Information**

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<thead>
<tr>
<th>Name</th>
<th>BioXF Name</th>
<th>Formula</th>
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**Formula Summary**

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**Formula Details**

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**Compound Spectrum**

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**Figure 7**

Detailed metabolite report about the formula, including isotopic pattern, calculated for dihydroxy metabolite of buspirone at retention time 0.71 min.

**Figure 8**

Detailed metabolite report for dihydroxy metabolite buspirone at retention time 0.71 min:
A) Extracted Ion Chromatograms (EIC) of compounds with mass 418.24
B) Extracted Compound Chromatogram (ECC) of dihydroxy metabolite of buspirone at retention time 0.71 min
C) Measured and calculated isotopic pattern of dihydroxy buspirone metabolite at retention time 0.71 min.
**Conclusion**

This Application Note demonstrated the use of the Agilent 1290 Infinity LC system with an Agilent Q-TOF LC/MS system for fast separation and accurate mass measurement of compounds in an in-vitro metabolite sample under high-throughput conditions. The metabolite compounds were separated in a run time below one minute and the width of the peaks extracted by the Metabolite ID software were below one second (FWHH). The major metabolites were identified quickly by means of the Agilent Metabolite Identification software. A summary report as well as detailed reports for each metabolite were generated.

**References**
