

Analysis Report

Bruker IVDr **Quant**ification in **UR**ine B.I.Quant-UR b^{TM}

Sample ID: ALZ_Urine_Rack01_RCM_221214_expno790.100000.10r

Measuring Date: 24-Dec-2014 08:30:32

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Quantification Method Version: Quant-UR B.1.1.0

Disclaimer

RESEARCH USE ONLY: This is no clinical diagnostic analysis report. Must not be used for clinical (medical or IVD) diagnosis or for patient management! Additional concentration range information (95% range) provided numerically or graphically in this report must not be used for clinical diagnostic interpretation.

Application of B.I.Quant-UR B 1.1.0 requires use of Bruker's B.I.Methods SOP for urine.

Summary

The following metabolites were found with concentrations outside the 95% range of Bruker Quant-UR B.1.1.0 urine metabolite concentration database:

<u>Amino acids and derivatives:</u> 4-Aminobutyric acid (43 mmol/mol Crea), Betaine (< 7 mmol/mol Crea), Carboxylic acids: Acetic acid (80 mmol/mol Crea),

<u>Keto acids and derivatives:</u> Acetoacetic acid (34 mmol/mol Crea), Oxaloacetic acid (81 mmol/mol Crea).

Further detailed information is provided on the following pages.



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1 Creatinine

Compound	Conc.	LOD	r	ρ	Δ	95% Range ^(*)
	mmol/L	mmol/L	mmol/L	%	mmol/L	mmol/L
Creatinine	9.2	0.3	9.170	100	0.106	1 - 19

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

2 Amines and derivatives

Compound	Conc.	Conc.	LOD	r	ρ	Δ	95% Range ^(*)
	mmol/L	_mmol_ mol Crea	_mmol_ mol Crea	mmol/L	%	mmol/L	<u>mmol</u> mol Crea
Dimethylamine	0.31	34	31	0.313	100	0.012	≤ 54 🔲
Trimethylamine	< 0.02	< 2	2	0.006	80 🔾	0.002	≤ 3 □ □

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

3 Amino acids and derivatives

Compound	Conc.	Conc.	LOD	r	ρ	Δ	95% Range ^(*)
	mmol/L	mmol mol Crea	mmol mol Crea	mmol/L	%	mmol/L	mmol mol Crea
1-Methylhistidine	< 0.14	< 15	15	0.000	00	0.809	≤ 15 □
2-Furoylglycine	< 0.35	< 39	39	0.000	0 🔾	0.171	≤ 40 □
4-Aminobutyric acid	0.39	43	20	0.392	66 🔾	0.566	≤ 20 □ □
Alanine	0.23	25	10	0.225	99 🔵	0.024	11 - 72
Arginine	< 6.9	< 750	750	0.973	0 🔾	2.062	≤ 750 □ □
Betaine	< 0.06	< 7	7	0.055	100	0.009	9 - 78
Creatine	0.60	65	50	0.600	100	0.106	≤ 280 🔲
Glycine	0.74	80	34	0.737	100	0.031	38 - 440
Guanidinoacetic acid	< 0.95	< 100	100	0.419	99 🔵	0.051	≤ 140 🔲
Methionine	< 0.16	< 18	18	0.000	0 🔾	0.331	≤ 18 <u> </u>
N,N-Dimethylglycine	0.06	7	5	0.064	99 🔵	0.047	≤ 15 🔲
Sarcosine	< 0.02	< 2	2	0.003	0 🔾	0.014	≤ 7 □□□
Taurine	< 1.3	< 140	140	0.129	80 🔾	0.166	≤ 170 🗔
Valine	< 0.02	< 2	2	0.009	88 🔵	0.015	≤ 7 □□□

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.



4 Benzene and substituted derivatives

Compound	Conc.	Conc.	LOD	\mathbf{r}	ρ	Δ	95% Range ^(*)
	mmol/L	_mmol_ mol Crea	_mmol_ mol Crea	mmol/L	%	mmol/L	_mmol_ mol Crea
Benzoic acid	< 0.09	< 10	10	0.000	0 🔾	0.072	≤ 10 □
D-Mandelic acid	< 0.02	< 2	2	0.000	0 🔾	0.223	2 - 17
Hippuric acid	2.0	210	170	1.965	99	0.187	≤ 660 🔟

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

5 Carboxylic acids

Compound	Conc.	Conc.	LOD	r	ρ	Δ	95% Range(*)
	mmol/L	mmol mol Crea	mmol mol Crea	mmol/L	%	mmol/L	mmol mol Crea
Acetic acid	0.73	80	5	0.734	100	0.041	≤ 51 □ □ □
Citric acid	2.6	280	40	2.596	100	0.335	≤ 700 🔟
Formic acid	< 0.09	< 10	10	0.029	83 🔾	0.011	≤ 43 □ □
Fumaric acid	< 0.02	< 2	2	0.007	97 🔵	0.001	≤ 3 □ □
Imidazole	< 0.44	< 48	48	0.076	75 🔾	0.053	≤ 48 🔲
Lactic acid	< 0.44	< 49	49	0.377	99 🔵	0.059	≤ 110 🗔
Proline betaine	0.29	31	25	0.286	88 🔾	0.057	≤ 280 🔲
Succinic acid	0.05	6	5	0.051	90 🔾	0.015	≤ 39 🔃
Tartaric acid	0.10	11	5	0.099	99 🔵	0.009	≤ 110 🔲
Trigonelline	< 0.32	< 35	35	0.205	100	0.005	≤ 67 □□□

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

6 Fatty acids and derivatives

Compound	Conc.	Conc.	LOD	r	ρ	Δ	95% Range ^(*)
	mmol/L	mmol mol Crea	mmol mol Crea	mmol/L	%	mmol/L	_mmol mol Crea
2-Methylsuccinic acid	< 0.44	< 48	48	0.000	0 🔾	0.259	≤ 48 □ □

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.



7 Keto acids and derivatives

Compound	Conc.	Conc.	LOD	\mathbf{r}	ρ	Δ	95% Range ^(*)
	mmol/L	_mmol_ mol Crea	_mmol_ mol Crea	mmol/L	%	mmol/L	_mmol_ mol Crea
2-Oxoglutaric acid	< 0.85	< 92	92	0.129	27 🔾	0.253	≤ 92 □ □
3-Hydroxybutyric acid	< 0.95	< 100	100	0.000	0 🔾	1.775	≤ 100 🔲
Acetoacetic acid	0.31	34	14	0.307	90 🔵	0.117	≤ 30 □ □
Acetone	0.02	2	2	0.021	90 🔵	0.008	≤ 7 🔟
Oxaloacetic acid	0.75	81	17	0.746	97 🔵	0.230	≤ 66 □ Ⅰ
Pyruvic acid	< 0.08	< 9	9	0.061	99 🔵	0.006	≤ 13 🔲

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

8 Purine, Pyridine and Pyrimidine derivatives

Compound	Conc.	Conc.	LOD	r	ρ	Δ	95% Range ^(*)
	mmol/L	_mmol_ mol Crea	mmol_ mol Crea	mmol/L	%	mmol/L	_mmol mol Crea
1-Methyladenosine	< 0.04	< 5	5	0.000	00	0.168	≤ 5 □
1-Methylnicotinamide	< 0.29	< 32	32	0.043	98	0.006	≤ 32 □ □
Adenosine	< 3.6	< 390	390	0.000	0 🔾	2.211	≤ 390 □ □
Allantoin	0.18	20	17	0.180	97 🔵	0.026	≤ 47 🔟
Allopurinol	0.10	11	10	0.097	92 🔾	0.054	≤ 11 □
Caffeine	< 0.42	< 45	45	0.222	96	0.204	≤ 61 □
Inosine	< 0.17	< 19	19	0.032	25 🔾	0.087	≤ 19 🗔

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.

9 Sugars and derivatives

Compound	Conc.	Conc.	LOD	\mathbf{r}	ρ	Δ	95% Range ^(*)
	mmol/L	mmol_ mol Crea	mmol mol Crea	mmol/L	%	mmol/L	_mmol mol Crea
D-Galactose	< 0.40	< 43	43	0.000	00	0.049	< 44 □ □ □
D-Glucose	0.39	42	34	0.387	90 🔵	0.087	≤ 140 🔲
D-Lactose	< 0.88	< 96	96	0.121	28 🔾	0.115	≤ 96 □ □
D-Mannitol	< 1.7	< 180	180	0.000	0 🔾	5.301	≤ 180 🔲
D-Mannose	< 0.05	< 6	6	0.000	0 🔾	0.093	≤ 8 □ □
Myo-Inositol	< 41	< 4400	4400	0.000	0 🔾	8.428	≤ 4400 □

^(*) Gray horizontal boxes represent 95% concentration range, black vertical lines represent sample value.



10 Explanations

This section contains the definition of the parameters used above. In the section 10.1 a short manual, how to interpret the results, is presented. The section 10.3 contains the exact definitions of the parameters \mathbf{r} , ρ and Δ .

10.1 How to read the result

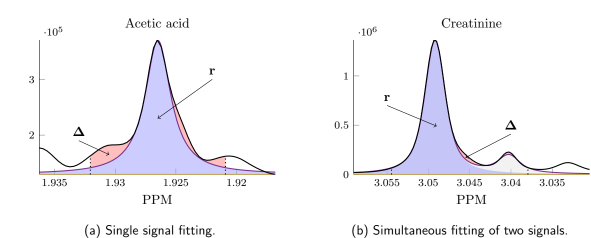


Figure 1: Examples of fitting.

In the figure 1(a), the black line, the blue line and the yellow line represent the original spectrum, the calculated signal fit and its baseline, respectively.

The blue area relates to the metabolite concentration to be determined and the red area represents a residue.

In case of the signal overlap a different approach is used: two or more overlapping signals are being fitted simultaneously. The most iconic example of such signals are the ones generated by CH_3 groups of Creatinine and Creatine. In such a case, the blue line and the grey area relate the sum of all fitted signals. The blue area corresponds to the concentration of the metabolite of interest (cf. figure 1(b)).

10.2 Result parameters

- a) Conc. is the final result concentration of the metabolite,
- b) **LOD** is the *limit of detection* of the given metabolite,
- c) ${\bf r}$ is the *raw concentration* i.e. the concentration equivalent of the resulting signal fit prior to comparing to **LOD** (relates to the blue area, cf. α)),
- d) ρ is the correlation of lineshape metabolite signal with calculated fit characterizing the match between metabolite signal and fit (cf. β)). Depending on the value of ρ , the following flag is displayed:



- , if the correlation is 95%,
- O, if the correlation is in between 85% and 95%,
- (), if the correlation is less than 85%,
- e) Δ is the concentration equivalent of the difference between metabolite signal and calculated fit (residue corresponding to the the red area, cf. γ)).

10.3 Detailed definitions

Let s, f and b denote the functions describing the *raw spectra*, *fitted curve* and *(fitted) baseline* respectively. These functions are chosen such that $s \approx f + b$. Moreover, let I be a relevant PPM interval and P_N be the proton number for given metabolite/signal.

 α) **r** (*raw concentration*) is defined as

$$\mathbf{r} = \frac{1}{P_N} \int_{\mathbb{R}} f(\xi) \, \mathrm{d}\xi.$$

 β) ρ is the *correlation* of the functions s and f+b, i.e.

$$\rho = \max(0, \operatorname{corr}(\overline{s}, \overline{f+b})),$$

where \overline{s} , $\overline{f+b}$ are numerical representations of the functions s and f+b on sufficiently fine mesh of the interval I.

 γ) Δ is the the area between the raw signal s and the fitted data f+b on the interval I expressed in the terms of the concentration, i.e.

$$\mathbf{\Delta} = \frac{1}{P_N} \int_I |s(\xi) - f(\xi) - b(\xi)| \, d\xi.$$