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Background. Metabolomics is a relatively new diagnostic tool that allows a deep insight into the body metabolism at a cellular level.

Objective. This paper provides a comprehensive view into the metabolomics methodology and shows usefulness of this approach in diagnosing and stratifying lung and breast cancers.


Results. In general, the metabolomic approach comprises three steps: 1) sampling and preparing biofluids or tissue homogenates, 2) identification of low-molecular weight compounds up to 1.0 kDa using nuclear magnetic resonance, mostly 1H-NMR and/or mass spectrometry, and finally 3) data processing and analysing. It is possible to identify a set of metabolites, which is specific for a certain metabolic status (the metabolic fingerprint). Furthermore, this set of metabolites provides information of possible pathomechanisms involved in the disease process i.e. information about the disease etiology. It has been proven that the change in metabolome precedes; not only clinical symptoms but other laboratory findings as well. Consequently, this approach, if sufficiently validated, seems to be very promising especially in screening and early diagnosing.

Conclusions. It was demonstrated that metabolomic approach allows to discriminate patients with cancer from healthy persons, as well as to differentiate between clinical stages of the cancer.

Key words: metabolomics; metabolome; breast cancer; lung cancer.

Introduction

To understand the complex processes occurring in the living systems the holistic approach should be used within employing all established multilevel approaches, currently is called ‘omics science’. In general, there are more than 100 types of omics subjects, including sciences such as: genomics, transcriptomics, proteomics, and metabolomics. This approach includes all-important live processes from information storage transcription processes, protein production, up to enzymes actions transforming certain substrates to products maintaining the cell living processes. While genomics and proteomics have been successfully introduced to routine diagnostics, metabolomics is still on the runway despite its assessed potential [1].

Metabolomics refers to the analysis and interpretation of ongoing processes in living organisms at the end of life chain consisting of genomics-transcriptomics-proteomics-metabolomics science. Metabolomics might be strongly influenced by external factors such as environment, life style, diet, medications, as well as by additional metabolite input from the body’s microbiomes. Metabolites are delineated as low-molecular weight compounds (LMWC) up to 1.0 kDa. Those compounds are carbohydrates, fatty acids, lipids, amino acids, nucleosides, or other organic molecules which are involved in biochemical reactions as substrates, intermediates, and/or final products. The set of body’s metabolites ultimately form its characteristic metabolome. Metabolome is a dynamic system, which is largely dependent on internal biochemical reaction and external factors mentioned above [1, 2]. Metabolomics is a comparative science, e.g. the obtained information should be compared to the referenced one or vice versa. With this approach metabolomics can be
regarded as a diagnostic tool, which in principle enables to describe current metabolic status of cells, tissues, and organs. Simply put, it can define the general health of a living organism. This is done by analysing body fluids, either secreted like urine, saliva, mother milk, stool, or obtained during standard medical sampling procedures such as serum, plasma, cerebrospinal fluid, tissue samples etc.. The samples collection for metabolomics studies is relatively simple, however certain procedures must be followed [3]. The patients must follow standard requirements: they should be fasting and reports all designated therapy, addictions, and medical history. The samples taken should be stored at -80 °C or preserved in liquid nitrogen [2, 4]. This low storage temperature ensures the maintaining of metabolic composition, i.e. minimizing possible changes in the metabolic profile after sampling, and protects against potential influence of the donor’s microbiome [5, 6].

Comprehensive or even partial determination of the metabolome is not possible with a single analytical method due to different levels of detection limits and different group determination. There are many analytical methods employed for metabolomics, each of them having both advantages and disadvantages. However, there are two main methods, which are routinely used in metabolomics studies: NMR (Nuclear Magnetic Resonance, mostly ¹H-NMR) and MS (Mass Spectrometry) hyphenated with separations techniques [2]. Studies based on both NMR and MS have a common pathway:

One of the most important advantages of the MS method is the sensitivity to femtomole concentration, which allows analysing up to several thousand molecules in a certain sample, resulting in precise molecular description in the tested specimen. On the other hand, reproducibility with MS is usually worse as compared to NMR spectroscopy. Unlike the MS, the NMR method provides high reproducibility with the ability to combine structure analysis and detection of the range of 30-90 molecules without sample destruction (biofluids). However, the detection limit of individual compounds is shifted several rows and strongly depend on the time of measurements (acquisition). The advantage of this method is the ability to measure different compounds such as amino acids, nucleosides, amines, acids, etc. all at the same time. The data obtained with both methods are both quantitative and qualitative among the set of detected metabolites [7].

The results obtained are further elaborated using chemometric multidimensional discriminatory methods. For this type of studies, the methods most commonly used are: unsupervised PCA, supervised PLS (Partial Least Squares Discriminant Analysis), and OPLS-DA (orthogonal version of PLS-DA). These approaches allow describing each subject not with 2 or 3 variables (chemical compounds or characteristic parameters) as in routine statistical analysis, but even hundreds of variables (chemical compounds - metabolites) can be analysed with selection of the most specific ones. Those compounds or more often sets of compounds can be employed as biomarkers for discriminating patients from healthy individuals or for staging of the disease [8].

Application of metabolomics in the diagnosis of cancer – selected studies

Lung cancer

In Europe, lung cancer accounts for approximately 20 % of all cancer deaths, with 376 000 deaths in 2008. Unfortunately, the early stages of this cancer are usually low-symptomatic and final diagnosis is made late in the natural history of the disease. Consequently, effective treatment is not possible up to 90 % of cases, and the overall 5-year survival rate is 11.2 % for men and 13.9 % for women [9]. The metabolomic studies of lung cancer are based on various body fluids, predominantly blood serum, urine, and saliva. The most
common type of lung cancer is non-small cell lung cancer (NSCLC). The main recognized risk factor for NSCLC is cigarette smoking. Additionally, COPD is considered an independent risk factor of lung cancer [10]. Our team has demonstrated a possibility to differentiate between patients with COPD and with two stages of NSCLC: an early NSCLC (E-NSCLC) and an advanced NSCLC (A-NSCLC). Using 1H NMR method we were able to identify 45 metabolites in patients’ serum. Concentrations of acetate, citrate, and methanol were significantly reduced in lung cancer subjects. In contrast, concentrations of N-acetylated glycoproteins, leucine, lysine, mannose, choline, lipids (L3 + L4), and two other unknown compounds were increased in for patients suffering cancer in all three performed comparisons: NSCLC vs. COPD, E NSCLC vs. COPD and ANSCLC vs. COPD [10]. Similar results were obtained by Musharraf et al [11]. Using GC-MS method for tracing metabolites in plasma, they were able to discriminate lung cancer patients from COPD patients, as well as healthy non-smokers and healthy smokers with high sensitivity (96.2 %) and specificity (92.05 %).

Another research group focused on differentiating early lung cancer patients from healthy controls. They used 1HMR and rapid resolution liquid chromatography (RRLC) methods to investigate metabolites in serum and identified 25 metabolites, which were up or down regulated. Those findings proved disorders in glycolysis, lipid metabolism, choline phospholipid metabolism, one-carbon metabolism, and amino acid metabolism. The use of both methods enabled diagnosing early stage of lung cancer with a very high accuracy [12]. Mass spectrometry hyphenated with gas chromatography (GC-MS) was used by Horia and co-workers [13]. They demonstrated that the levels of 23 of 58 serum metabolites and 48 of 71 sampled from tissue were significantly changed in patients with lung cancer with I-IV stage as compared, with healthy volunteers. An early cancer stage was also investigated by other group using LC-MS method. In this study, the achieved AUC value based on 12 metabolites was 0.836 [14].

Urine also has a strong discriminative potential in lung cancer diagnosing. This was proven in a study performed using 1H NMR method by Carrola and colleagues [15]. The main metabolites differentiating between healthy controls and lung cancer patients were: hippurate, trigonelline, β-hydroxyisovalerate, α-hydroxyisobutyrate, N-acetylglutamine, and creatinine. They were able to develop a classification model, which confirmed 93 % sensitivity and 94 % specificity with overall classification rate of 93.5 %. [15]. The studies performed by LC-MS method on urine samples from cohort of 469 patients with lung cancer and 536 controls revealed two biomarkers: creatine riboside and N-acetylneuraminic acid, which were significantly increased in non-small cell lung cancer and were associated with worse prognosis [16].

Exhaled breath condensate (EBC) seems to be naturally associated with lung cancer. Indeed, studies on EBC performed with GC-MS [17] and LC-MS [18] proved diagnostic usefulness of EBC differentiating lung cancer patients, COPD patients, smokers, ex-smokers, and healthy controls. In addition to the above mentioned biofluids, also sweat should be included, which was successfully used to differentiate the lung cancer patients from control individuals with risk factors and without them by its analysis by LC-MS system [19].

Recently, a very promising study has been published by Shen et al.. Using two-stage study design and advanced metabolon platform they were able to identify four metabolites, which may be useful biomarker candidates for identifying patients, who may benefit from platinum-based chemotherapy in advanced NSCLC [20]. The comprehensive review on lung cancer biomarkers and metabolomics methods has been published recently [21].

Breast cancer

Breast cancer (BC) is the most common cancer in women worldwide and the morbidity keeps rising. Incidence vary from 19.3 per 100,000 women in Eastern Africa to 89.7 per 100,000 in Western Europe [22, 23]. Some decrease in the number of death caused by BC is observed mainly due to the improvement in early diagnostics [24]. Nevertheless, routine methods currently used to detect BC usually are not effective, especially in the very early stages [25].

A new approach in cancer diagnostics involves the use of metabolomics tools for blood serum and urine analysis [25, 26, 27]. Metabolomic-based diagnosing not only seems to allow discriminating healthy controls from BC subjects, but also differentiating metastatic BC from early stages of BC as well [28, 29].

Previous studies proved that BC can be diagnosed by NMR spectroscopy and by MS, both by use of serum and urine [27, 30, 31].
However, those metabolomic studies involved relatively small groups and, utilized various protocols and methods [32]. Therefore, a more integrated and coherent methodology should be used [33].

The GC-MS method can be used for metabolic profiling of serum. Using this method, it was possible to differentiate patients with BC from patients with non-malignant tumours, and from healthy controls. Sets of amino acids, fatty acids, and lysolipids allowed differentiating these three groups [31]. There is also evidence that polar compounds analysis has been successfully applied for metabolomic-based BC diagnosing in tissue samples. Chae and co-workers used HR-MAS 1H NMR method for retrospective analysis in the patients with ductal carcinoma in situ (DCIS) diagnosed on preoperative biopsy. The univariate analysis proved that choline-containing compounds did not differ between the groups, while GPC/PC ratio, myo-inositol, and succinate were higher in the ‘pure’ DCIS group as compared to the invasive carcinoma subjects. Multivariate analysis OPLS-DA could discriminate to some degree between these two groups [34]. Comparable results were obtained in other groups. In a study based on add space 1H NMR method the authors were able to distinguish between the non-invasive intraductal carcinoma and invasive ductal carcinoma patients. Histidine concentration was significantly lower in the patients with invasive ductal carcinoma. In contrast, those patients presented higher concentrations of glucose, lactate, tyrosine, and lipids in plasma samples, as compared to the non-invasive carcinoma group [35].

It is also possible to differentiate metastatic BC patients from the ones with a localized, early-stage disease. Serum analysis with NMR spectroscopy identified 9 metabolites: histidine, acetacetate, glycerol, pyruvate, glycproteins (N-acetyl), mannose, glutamate, and phenylalanine concentrations, all of which were significantly different, than those in the other studied groups [26].

Volatile organic compounds (VOC) from urine samples can also be used for BC diagnosis. In the study of Silva et al., gas chromatography-mass spectrometry was used to obtain metabolomic patterns of 26 BC patients and 21 healthy individuals. Of the 79 volatile identified metabolites six compounds were of diagnostic power: (−)-4-carene, 3-heptanone, 1,2,4-tri-methylbenzene, 2-methoxythiophene, phenol, and dimethylsulfide. All of them were able to successfully discriminate between the groups [30].

More holistic approach was suggested by Bro et al. In a retrospective study they combined 1H NMR data with other relevant biological and phenotypic information to construct a patient’s biocountour. With this approach the authors could predict an increased risk of BC, a few years before its occurrence with sensitivity and specificity well above 80 % [36]. A very interesting paper aiming on possible associations of diet-related metabolites with BC risk was published recently by Pleydon et al.. Using CE-MS method they identified a bunch of metabolites, which were moderately correlated with increased risk of estrogen receptor related to BC development [37]. The annually increasing literature data proves the usefulness of metabolomics in BC [33, 38].

Conclusions

Data from the literature clearly demonstrate the usefulness of metabolomic approach in diagnostics of lung cancer, breast cancer, and many other diseases. Despite this, the metabolomic approach is still not routinely implemented into medical protocols. This results from many factors. Analytical methods used for metabolomic research are neither standardized nor validated. Consequently, it is difficult to: compare results obtained by various research groups, and determine the definitive clinical recommendations. Therefore we need more and more research to overcome these difficulties. However, in the future the metabolomic approach with fingerprinting and profiling-based methods, combined together with predictive-discrimination statistical models should be the method of choice for preventive, screening, and treatment research.

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