Review

Defining Acute Coronary Syndrome through Metabolomics

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Abstract: As an emerging platform technology, metabolomics offers new insights into the pathomechanisms associated with complex disease conditions, including cardiovascular diseases. It also facilitates assessing the risk of developing the disease before its clinical manifestation. For this reason, metabolomics is of growing interest for understanding the pathogenesis of acute coronary syndromes (ACS), finding new biomarkers of ACS, and its associated risk management. Metabolomics-based studies in ACS have already demonstrated immense potential for biomarker discovery and mechanistic insights by identifying metabolomic signatures (e.g., branched-chain amino acids, acylcarnitines, lysophosphatidylcholines) associated with disease progression. Herein, we discuss the various metabolomics approaches and the challenges involved in metabolic profiling, focusing on ACS. Special attention has been paid to the clinical studies of metabolomics and lipidomics in ACS, with an emphasis on ischemia/reperfusion injury.

Keywords: metabolomics; lipidomics; acute coronary syndrome; ischemia/reperfusion injury

1. Introduction

Coronary artery disease (CAD) continues to be a major public health concern with considerable morbidity and mortality in developed countries [1]. CAD includes chronic coronary artery disease (stable angina) and acute coronary syndrome (ACS), which almost invariably presents chest discomfort with or without dyspnea. It is estimated that in the United States alone, 720,000 people experience a new episode of ACS every year [2]. The costs and resource utilization associated with ACS also place an enormous economic burden on the healthcare system [2–4]. For instance, in a retrospective single-cohort study, it was found that the average one-year cost per patient associated with ACS was around 32,000 US dollars [4]. The term ACS refers to a spectrum of conditions in which myocardial ischemia or infarction develops due to acute occlusion of coronary blood flow to any part of the heart. The usual cause of acute occlusion is coronary artery thrombosis caused by rupture or erosion of a high-risk, lipid-laden, atheromatous plaque [5]. This sudden, reduced blood flow to the heart results in an imbalance between myocardial metabolic demands and blood supply, leading to myocardial ischemia, which is the hallmark of ACS [3,6]. The imbalance may also be caused by several other cardiac abnormalities, including...
coronary artery embolism, coronary spasm, coronary dissection, severe anemia, and calcific aortic valve stenosis [3]. Depending upon the range of ischemic state, location of the occlusion, cardiac biomarker levels (e.g., troponin), and ST-segment elevation on the electrocardiogram (ECG), ACS is mainly categorized into three types (Figure 1), namely unstable angina (UA), non-ST-segment elevation myocardial infarction (NSTEMI), and ST-segment elevation myocardial infarction (STEMI) [5]. Typically, a complete coronary artery occlusion leading to myocardial tissue injury and elevated cardiac troponin level results in STEMI. Partial occlusion or occlusion with collateral circulation may lead to NSTEMI or UA, depending on the presence or absence of cardiac troponin level, respectively [5,7]. Troponins are released into the bloodstream when there is any damage to the heart muscle [8]. Due to its high sensitivity and specificity, cardiac troponin measurement is an essential element in diagnosing and managing ACS.

Figure 1. Classification of acute coronary syndromes. Acute coronary syndromes are categorized into unstable angina (UA), non-ST-segment elevation myocardial infarction (NSTEMI), and ST-segment elevation myocardial infarction (STEMI). A complete coronary artery occlusion due to thrombus formation results in STEMI, where the coronary blood flow is completely obstructed. A partial occlusion of the artery (blood flow is not entirely restricted) can result in NSTEMI or UA.

The human heart is the most metabolically active organ, and under normal conditions, cardiac metabolism is tightly regulated by oxygen availability, substrate oxidation, hormonal and neurohumoral signals, among other factors [9]. The primary substrates for cardiac ATP production are lipids, carbohydrates, lactate, and glycogen [9]. Of these, lipids alone contribute to 60–90% of ATP production in the heart via oxidation of fatty acids [10]. Due to its exuberant use in the myocardium and its role in maintaining the myocardial cell structure and cardiac function, alterations in lipid metabolism contribute to many cardiovascular pathologies, including atherosclerosis, insulin resistance, hypertension, and type 2 diabetes mellitus [11]. A fall in oxygen level and substrate availability during myocardial ischemic conditions alters the dynamic equilibrium state of cardiac metabolism and leads to loss of homeostasis [12]. Hence, it is not surprising that ACS involves changes in cardiac metabolism. While some of the metabolic changes help the heart adjust to the altering substrate needs and physiological demands, most changes are maladaptive.
and initiate other cardiac abnormalities, including myocardial stunning, cell death, and contractile dysfunction [13]. Recent findings suggest that alterations in the cardiac metabolism alone can also perpetuate disturbances in systemic metabolism [14]. Hence, blood-derived plasma or serum metabolic profile could provide insights into the pathophysiological processes happening within the heart in the event of an ACS.

Metabolomics is the new entrant in the 'omics' cascade in the systems biology (genomics, transcriptomics, and proteomics) approach (Figure 2). It involves high-throughput identification and quantification of small chemical compounds (<1000 to 1500 Da), commonly known as metabolites, present in a variety of biological system such as a cell, an organism, or biological fluids [15,16]. The complete set of these small-molecule metabolites and their interactions within the biological system are known as the metabolome. The metabolome includes endogenous (e.g., amino acids, fatty acids, sugars, carbohydrates, vitamins, lipids, and their derivatives) as well as exogenous (e.g., pollutants, pharmaceuticals, food additives, xenobiotics) compounds. Lipidomics is a novel subdivision of metabolomics, dedicated to the detailed analysis of complex lipid mixtures found in biological materials [17]. The metabolome composition is inherently dynamic and flexible due to its continuous interaction within the biological system and also with the outside environment, including the effects of drugs, nutrition, lifestyle, or therapeutics [18]. Hence, any perturbations in metabolite levels are a true reflection of the phenotype and function of the developmental or pathological state of the biological system. Recent advancements in 'omics' technology platforms, particularly genomics and proteomics technologies, have enabled us to assess the changes within the genome and the proteome in cardiovascular diseases (CVD), including ACS. Metabolomics and lipidomics are required to bridge the knowledge gap between phenotype and metabolic abnormalities in CVD.

Given that ACS is a multifactorial phenomenon, it requires a global assessment of molecular/cellular pathological functions from both untargeted and targeted approaches. Recognizing this, the American Heart Association recently released a scientific statement highlighting the potential impact of metabolomics in cardiovascular health and disease [19]. Thus, metabolomics and lipidomics are now evolving as essential tools in cardiovascular research (i) to gain a better understanding of the mechanisms underlying CVD, (ii) to determine the clinically relevant metabolic perturbations, and (iii) to identify novel biomarkers involved in various cardiovascular disease conditions.
2. Analytical Tools in Metabolomics

Though metabolomics research is considered a relatively new field, the initial report of screening metabolites in body fluids for diagnostic purposes can be traced back to ancient China (1500–2000 BC), where traditional Chinese physicians used ants to diagnose diabetes by evaluating the smell, taste, and color of urine samples from patients [20]. However, the pioneering work carried out by the laboratories of Robinson’s and Pauling’s research groups in the early 1970s [21–23] laid the foundation for a new era in metabolomics research. They successfully quantitated metabolites from various biofluids, including human blood and urine vapor, by applying the gas chromatography coupled with mass spectrometry (GC/MS) technique. The first printed reference to the word ‘metabolic profile’ appeared in the literature in 1971 [24,25], when Horning and Horning used the GC/MS platform to profile different human metabolic products, including sugars, alcohols, and drug metabolites from blood and urine samples. Since these initial efforts, several investigators have contributed in parallel to develop this naive concept into a format that is on par with the other established ‘omics’ approaches, namely genomics and proteomics. The word ‘metabolome’ was first used by Oliver Fiehn, a key metabolomics researcher, in 1998 [26] to denote the changes in relative concentrations of metabolites due to the deletion or overexpression of a gene. Due to the metabolome’s vast complexity and dynamic nature, no single instrument platform currently available can simultaneously analyze all the metabolites present in a biological sample. For example, within lipids alone, diverse lipid classes and lipid molecular species are categorized based on their structural and chemical properties. Moreover, while some lipids such as cholesteryl esters and triglycerides are usually present in high concentrations (1–10 mmol/L) in samples such as plasma, others such as eicosanoids derived from arachidonic acid are present only in trace amounts.
amounts (1–10 pmol/L) [27]. Numerous analytical platforms with varying degrees of specificity and sensitivity have been developed in the last two decades to measure specific portions of the complex metabolome. Presently, nuclear magnetic resonance (NMR) spectroscopy and mass spectrometry (MS), coupled with gas or liquid chromatography (GC/MS or LC/MS, respectively), are the most widely used analytical tools in metabolomics investigations [15]. In addition, analytical platforms such as MS coupled with capillary electrophoresis (CE/MS), shotgun lipidomics, and direct MS infusion methods are also used to fully understand the broad spectrum of metabolites [15,28]. For instance, while NMR makes the study of different isomers possible, GC/MS or LC/MS can analyze a large variety of molecules with high sensitivity, and CE/MS can provide data of high resolution. Table 1 summarizes the main advantages and disadvantages of using MS and NMR techniques.

| Technique | Advantages | Disadvantages |
|-----------|------------|---------------|
| NMR       | Highly reproducible results | Relatively low sensitivity compared to MS |
|           | Provides structural information about the compounds | Only suited for medium to high abundant metabolites (micro-molar range) |
|           | Minimal requirement for sample preparation | Relatively longer data acquisition times compared to MS |
|           | Non-destructive in nature and suitable for multiple analyses of the same sample | Highly pH-sensitive |
|           | Allows investigation of tissue energetics and in vivo metabolism | |
|           | Suitable for compounds which are otherwise difficult to ionize or derivatize | |
|           | Appropriate to use with samples with high salt content, including urine | |
|           | Well-established NMR spectra library to aid data analysis | |
|           | Ability to detect different isomeric products | |
| GC/MS     | Method of choice for the analysis of volatile/non-polar metabolites | Detection of polar metabolites is difficult and needs chemical derivatization. |
|           | Increased signal-to-noise (S/N) ratios and relatively better resolution | Limited metabolome coverage |
|           | Publicly available spectral libraries for compound identification | The high temperature applied in GC/MS can cause degradation or transformation of compounds. |
| LC/MS     | Ability to analyze metabolites with a wide range of polarity, including thermally unstable ones | Not suitable for the analysis of gaseous mixtures |
|           | Quicker and less extensive sample extraction procedures | Decreased sensitivity due to ion suppression |
|           | Suitable for measurement of compounds of lower volatility | Difficulty in distinguishing isomers (both structural and positional) of molecules |
|           | Requires little sample volume | |

Abbreviations: NMR, nuclear magnetic resonance spectroscopy; GC/MS, gas chromatography/mass spectrometry; LC/MS, liquid chromatography/mass spectrometry.
2.1. NMR Spectroscopy

NMR spectroscopy is one of the most prevalent techniques used in metabolomics research for structural elucidation of both pure compounds and mixtures. This technique relies on the magnetic spin properties (nuclear magnetic moment and angular momentum) of protons and neutrons in the nucleus of an atom to provide information on molecular structures. It is based on the principle that any nucleus possessing a (non-zero) nuclear spin can release characteristic radiofrequency waves when placed in an external magnetic field. The resonance frequency of the released energy is affected by the chemical microenvironment of these atoms and the coupling effect with nearby nuclei. The resulting frequencies are recorded as chemical shifts, and their patterns can be used for compound identification [15]. The most commonly used nuclei for analysis are $^1$H, $^{13}$C, and $^{31}$P isotopes. The proton NMR spectroscopy ($^1$H-NMR) has the advantage of having greater relative sensitivity and a short acquisition time. For this reason, $^1$H-NMR is widely used to identify and quantitate a large number of small molecules coexisting in biological material, such as tissues, whole cells, and biofluids. The major disadvantage of traditional one-dimensional (1D) $^1$H-NMR is that the spectrum became very complex for larger, more complex molecules, making interpretation difficult as most of the signals overlap heavily. However, two-dimensional (2D) NMR helps circumvent this challenge by resolving signals that usually overlap in $^1$H-NMR. In short, 2D-NMR provides extra information about a molecule compared to traditional $^1$H-NMR spectra, thereby making spectral interpretation easy. Current practice requires 1D $^1$H-NMR to complement 2D-NMR or mass spectrometer for molecular identity confirmation. Carbon-13 NMR spectroscopy ($^{13}$C-NMR) and phosphorous-31 NMR spectroscopy ($^{31}$P-NMR) predominantly find applications in cellular energetics, particularly in tracking cellular changes in cardiac metabolism under normoxic and ischemic conditions [29–31].

Table 2 shows that in ACS metabolomics studies, $^1$H-NMR spectroscopy is the most used NMR technique for analyzing biofluids, including plasma, serum, and urine. $^1$H-NMR spectroscopy is routinely used to analyze several low-molecular-weight molecules, including branched-chain amino acids (e.g., valine, isoleucine) and ketone bodies from plasma and serum to find biomarkers for ACS [32–34]. In a study looking at potential candidate biomarkers for UA from plasma samples, $^1$H-NMR spectroscopy identified ten steroid metabolites belonging to the steroid hormone biosynthesis pathway, including one mineralocorticoid (deoxycorticosterone) and nine sex hormone metabolites (e.g., estradiol, 2-hydroxyestradiol) [35]. Additionally, NMR has become the primary tool for analyzing urinary metabolites. In a study involving UA patients [36], $^1$H-NMR analysis could successfully identify and quantitate waste metabolites such as trimethylamine-N-oxide (TMAO) and trimethylamine (TMA) from urine specimens, in addition to short-chain fatty acids (e.g., 3-hydroxybutyrate), organic acids (e.g., indol-3-acetate, methylmalonate), and amino acids (e.g., lysine, proline).

The main advantages of NMR are its non-destructive nature and higher reproducibility [37,38]. However, it is often limited by its requirement of a larger sample amount (2–50 mg) and therefore reduced sensitivity for low abundant compounds [39]. An early metabolomics study showed that 1H-NMR spectroscopy could correctly diagnose the presence of CAD and assess its severity [40]. Other applications of NMR-based metabolic profiling in ACS mainly include analysis of both urine [36] and serum [33,34] metabolites for unstable angina pectoris disease, investigating the serum metabolic characteristics of acute myocardial infarction (AMI) patients in comparison with those of chest pain controls [41], and deciphering the metabolic fingerprint of coronary blood in STEMI patients [42].
| No. | First Author, Year | Sample Size | Sampling Time | Specimen/Technique | Main Findings |
|-----|-------------------|-------------|---------------|--------------------|---------------|
| 1   | W Zhong [43], 2021 | 284 ACS; 130 HC | At the time of hospital admission | Plasma LC/MS | Phenylalanine, arginine, and proline metabolism and synthesis and degradation of ketone bodies are involved in ACS pathogenesis. |
| 2   | E Chorell [44], 2021 | 50 STEMI; 50 NSTEMI; 100 HC | After fasting for 4 h | Plasma GC/MS, LC/MS | Plasma lysophospholipids ratio (LPC:LPE) could predict future risk in STEMI and NSTEMI patients. |
| 3   | N Aa [45], 2021   | 85 MI; 61 non-MI chest pain; 84 HC | Within 6 h of the initial symptom attack | Plasma GC/MS, LC/MS | Patients with MI had elevated plasma levels of deoxyuridine, methionine, and homoserine. |
| 4   | H Chen [46], 2021  | Discovery: 942 Validation: 493 | After fasting for 8 h | Plasma LC/MS | Perturbations in cysteine and methionine metabolism and glycerophospholipid metabolism are associated with CAD severity. |
| 5   | A Mehta [47], 2020 | Discovery: 454 Validation: 322 | After overnight fasting | Plasma LC/MS | 23 differential metabolites were identified between AMI and NOCAD, including 12 acylcarnitines, 7 fatty acids, 3 glycerophospholipids, and L-tryptophan. |
| 6   | J Li [48], 2020    | 136 NOCAD; 118 AMI | After overnight fasting | Serum LC/MS | A total of four metabolites including isoundecylic acid, betaine, 1-heptadecanoyl-sn-glycero-3-phosphocholine, and acetylcarnitine could discriminate stable and vulnerable plaques. |
| 7   | H Jiang [49], 2020 | 252 ACS | After initial diagnosis of ACS | Serum LC/MS | L-homocysteine sulfonic acid, cysteic acid, and carnitine could serve as predictive markers for AMI risk. |
| 8   | A Khan [50], 2020  | 112 patients at AMI risk; 89 HC | During routine blood collection after overnight fasting | Serum LC/MS | 17 metabolites involved in pathways such as steroid hormone biosynthesis, aminoacyl-tRNA biosynthesis, and lysine degradation could serve as promising biomarkers for UA diagnosis. |
| 9   | M Pouralijan Amiri [35], 2020 | 94 UA; 32 controls (angina, but no CAD) | After coronary angiography | Plasma H-NMR | Coronary blood fingerprint in STEMI patients was represented by choline, phosphocholine, myo-inositol, lysine, ornithine, and 2-phosphoglycerate metabolites. |
| 10  | A Vignoli [32], 2019 | 825 total, 702 survivors and 123 deceased | 24–48 h after the PCI and overnight fasting | Serum H-NMR | Characterization of metabolite–metabolite association, can be used as a potential tool to predict mortality in AMI patients. |
| 11  | G Gundogdu [51], 2020 | 20 STEMI; 15 HC | Within an hour of the initial symptom attack | Serum LC/MS | Malonic acid, maleic acid, fumaric acid, and palmitic acid could be used for the diagnosis of STEMI. |
| 12  | A Surendran [52], 2019 | 27 STEMI | Pre-PCI, 2, 24, and 48 h post-PCI | Plasma LC/MS | Identified lipids and lipid-derived molecules as the major constituents of the altered metabolomic profile prior to PCI and in the follow-up time intervals post-PCI. |
| 13  | J Wang [53], 2019   | 40 UA; 39 HC | Blood samples taken at the same day of inclusion in the study | Plasma LC/MS | 27 metabolites, including free fatty acids, amino acids, LPE, LPC, and organic acids, can be used to diagnose UA patients. |
| 14  | M Deidda [42], 2019 | 15 STEMI | Coronary artery blood sampling during PCI | Plasma H-NMR | Coronary blood fingerprint in STEMI patients was represented by choline, phosphocholine, myo-inositol, lysine, ornithine, and 2-phosphoglycerate metabolites. |
| 15  | A Vignoli [54], 2019 | Training: 80 survivors and 40 deceased Validation: 752 survivors and 106 deceased | 24–48 h after the PCI and overnight fasting | Serum H-NMR | Mortality in AMI patients was associated with elevated serum levels of acetone, 3-hydroxybutyrate, mannose, creatinine, acetate, formate, proline, and lower serum levels of valine and histidine. |
| 16  | VAM Goulart [55], 2019 | 15 STEMI; 19 HC | Within 7 h after hospitalization | Plasma LC/MS | STEMI metabolic fingerprint includes perturbations associated with phosphatidylcholines, lysophosphatidylcholines, sphingomyelins, and biogenic amine species. |
| 17  | Y Wang [56], 2018   | 36 ACS; 30 HC | Not specified | Urine LC/MS | Identified fatty acid metabolism, fatty acid β-oxidation, amino acid metabolism, and TCA cycle as critical pathways associated with ACS pathogenesis. |
| No | Author | Year | Case Description | Sample | Analytical Approach | Findings |
|---|---|---|---|---|---|---|
| 18 | X Du [57], 2018 | 96 STEMI with post-PCI AEs; 96 without AEs | Arterial blood before coronary angiography | Plasma LC/MS | Circulating levels of branched-chain amino acids (BCAAs) were associated with the risk of adverse cardiovascular events in STEMI patients. |
| 19 | X Du [58], 2018 | 138 STEMI with AHF; 138 STEMI without AHF | At the time of hospital admission | Plasma LC/MS | Elevated plasma BCAA levels were associated with long-term adverse cardiovascular events in patients with STEMI and AHF. |
| 20 | L Huang [59], 2018 | 44 STEMI (22 LMCAD and 22 non-LMCAD); 22 HC | At the time of hospital admission | Plasma LC/MS | Retinol metabolism was the most perturbed metabolic pathway for the LMCAD phenotype. |
| 21 | D Dazhi [41], 2018 | 45 AMI; 45 chest pain controls (CPCS) | At the time of hospital admission and prior to any medication | Serum H-NMR | Multiple altered metabolic pathways, including the TCA cycle, lipoprotein changes, anaerobic glycolysis, gluconeogenesis, and fatty acid metabolism, characterize AMI patients compared to CPCS. |
| 22 | M Kohlhauer [60], 2018 | 115 STEMI; 26 control patients (SA/NSTEMI) | Immediately after stent deployment | Plasma LC/MS | Increased levels of myocardial succinate are found in STEMI patients. |
| 23 | L Zhang [61], 2018 | 2,324 patients who underwent coronary angiography | Before coronary angiography | Plasma LC/MS | N-acetyl neuraminic acid plays a key role during CAD progression. |
| 24 | X Yin [62], 2018 | 20 STEMI; 20 non-ACS patients | Pre-PCI | Plasma LC/MS, ICP/MS | ACS patients are characterized by disturbances in LPC, caffeine, glycolysis, tryptophan, and sphingomyelin metabolism. |
| 25 | W Yao [33], 2017 | 22 UA; 22 HC | Within 24 h after overnight fasting | Serum H-NMR | UA patients are characterized by perturbations in phospholipid and amino acid metabolism. |
| 26 | SE Ali [63], 2016 | 30 STEMI; 15 UA; 15 HC | 1–2 h post-angioplasty for UA patients | Serum GC/MS, SMPE-GC/MS, H-NMR | Elevated levels of serum hydrogen sulfide could discriminate STEMI patients from UA patients. |
| 27 | Y Fan [64], 2016 | Discovery: 1086 Validation: 933 | Before coronary angiography | Plasma GC/MS | 89 differential metabolites were identified between and within different CAD subtypes. |
| 28 | X Xu [65], 2015 | 38 SA; 34 AMI; 71 HC | After overnight fasting | Serum LC/MS | Different lipid classes, including fatty acids, steroids, phospholipids, sphingolipids, and glycerolipids, are associated with CAD progression. |
| 29 | L Huang [66], 2016 | 47 STEMI (23 youth, 24 elderly), 48 healthy controls (24 youth, 24 elderly) | Post-PCI | Plasma LC/MS | The most perturbed metabolic pathway in young STEMI patients was sphingolipid metabolism. |
| 30 | K Ameta [34], 2016 | 65 UA; 62 HC | Within 4 h of onset of angina | Serum H-NMR | Five significantly altered metabolites, namely valine, alanine, glutamine, 5-oxoproline, and adenine, differentiate UA patients from HC. |
| 31 | Z Li [36], 2015 | 27 UA; 20 HC | In the morning after fasting for 12 h | Urine H-NMR | 20 metabolites, including energy metabolism-related metabolites and amino acids, could discriminate UA patients from HC. |
| 32 | S Naz [67], 2015 | Discovery: 16 STEMI; 16 NSTEMI Validation: 20 STEMI; 28 NSTEMI | Pre-PCI | Serum LC/MS | Carnitine-related compounds and amino acids were differentially present in STEMI and NSTEMI conditions. |
| 33 | CM Laborde [68], 2013 | Discovery: 35 NSTEACS; 35 HC Validation: 15 NSTEACS; 15 HC | At the onset of the syndrome | Plasma GC/MS, LC/MS | A panel of metabolites consisting of 5-OH-tryptophan, 2-OH-butyric acid, and 3-OH-butyric acid could serve as markers for the early diagnosis of ACS. |
| 34 | M Sun [69], 2013 | 45 UA; 43 atherosclerosis controls | In the morning after overnight fasting | Plasma LC/MS | 16 potential endogenous biomarkers for UA were identified including kynurenine. |
| 35 | J Teul [70], 2011 | 19 NSTEACS; 6 HC | Immediately before coronary angiography, day 4, 2 months and 6 months after diagnosis | Plasma GC/MS | 27 metabolites including glucose, fructose, myoinositol, pyruvate, lactate, and succinate varied with time following an ACS event. |
| 36 | M Vallejo [71], 2009 | 9 NSTEACS; 10 stable in the morning after fasting atherosclerosis; 10 HC | Plasma GC/MS | Plasma fingerprinting characterizes a key role for 4-hydroxyproline in ACS. |

**Abbreviations:** STEMI, ST-elevation myocardial infarction; NSTEMI, non-ST-elevation myocardial infarction; NSTEACS, non-ST-elevation ACS; NOCAD, nonobstructive coronary artery disease; AMI, acute myocardial infarction; AHF, acute heart failure; UA, unstable angina; HC, healthy control; CPCS, Coronary Patient Control Study.
heart failure; MI, myocardial infarction; SA, stable angina pectoris; UA, unstable angina pectoris; PCI, percutaneous coronary intervention; ACS, acute coronary syndromes; CAD, coronary artery disease; HC, healthy control; LC/MS, liquid chromatography/mass spectrometry; GC/MS, gas chromatography/mass spectrometry; H-NMR, proton nuclear magnetic resonance; SPME, solid-phase microextraction; LPC, lysophosphatidylcholine; LPE, lysophosphatidylethanolamine; TCA, tricarboxylic acid cycle.

2.2. Mass Spectrometry

Mass spectrometry (MS) provides accurate weight measurements of one or more molecules within a sample of interest. MS separates the molecules based on their specific mass-to-charge ratio \((m/z)\) by converting them into ions in the gas phase [72]. The separated ions are then sorted according to their acceleration and deflection in an external electromagnetic field. The final output is presented as the relative abundance of each ion as an \(m/z\) spectrum. MS is extensively used in the field due to its wide dynamic range, speed, high sensitivity, and the ability to identify and quantify more metabolites in a single measurement relative to NMR.

The MS essentially has three main components: (1) the ion source, where the sample is ionized, (2) the mass analyzer, where the ions are separated according to their mass-to-charge \((m/z)\) values, and (3) the detector, which provides a count of the intensity of separated ions [73]. The most commonly used mass analyzers are Time-of-Flight (TOF), magnetic sector, and quadrupole, each with its own set of strengths and limitations [74]. Time-of-Flight, as the name implies, uses a flight tube of known length, where the ions are separated based on the flight time (time taken for the ions to travel through the flight tube). TOF MS consists of a pulsed ion source and therefore is best suited with ionization methods that ionize molecules in pulses, such as laser ionization. TOF systems have an excellent mass range and are generally utilized for high-resolution MS. Unlike TOF systems, magnetic sector mass analyzers use a magnetic field to sort ions of different mass-to-charge ratios. The key advantages of magnetic sector mass analyzers are their high sensitivity and high resolution. As its name suggests, quadrupole consists of four parallel cylindrical or hyperbolic rods. The opposite rods are connected electrically inside a vacuum chamber. By changing the electrical potential, ions with different \(m/z\) values can be ‘filtered’ through the quadrupole to the detector one after another. The quadrupole mass analyzers are usually compact and have good scan speed, durability, and reliability. However, they usually have a limited mass range.

MS uses various ionization methods for mass analysis. The classic methods include matrix-assisted laser desorption ionization (MALDI) and electrospray ionization (ESI), in addition to other methods, such as electron impact (EI) ionization, chemical ionization, and atmospheric pressure chemical ionization (APCI) [75]. ESI is based on the evaporation of charged droplets and is notable for being the softest ionization method, allowing for generating ions with multiple charges with great sensitivity and no matrix interference [76]. However, some of the pitfalls of ESI include incompatibility with salts, complex mixtures, and impure samples. On the other hand, MALDI functions by proton absorption and transfer and can ionize metabolites with much higher masses (up to 300,000 Da). MALDI is also compatible with both salts in millimolar concentration as well as complex samples. One disadvantage of using MALDI is the potential for the degeneration of metabolites due to the acidic matrix or photogeneration via laser ionization. Additionally, MALDI is also not suitable to analyze low-molecular-weight compounds (<1000 \(m/z\)) because of the matrix background interferences in this mass range [77]. Overall, it has been shown that using multiple ionization approaches with mass spectrometry can enhance the detection of global metabolome for complex samples [78].

A separation technique in a time dimension (GC, LC, or CE) is often employed before MS analysis to isolate the individual components from complex samples containing hundreds to thousands of small molecules [79]. The chromatographic separation techniques use the physicochemical properties of the compounds in a sample, such as polarity, size, and presence of double bonds, to separate the molecules inside the medium [80]. Hence,
combinations of separation and MS, usually either GC/MS or LC/MS, have become the preferred analytical choice for small-molecule analysis from complex biological samples. The application of MS-based metabolic fingerprinting in ACS mainly includes investigating biomarkers for ACS [68–71], characterizing the metabolic difference between different types of ACS [64,65,67], and identifying the molecular differences between patients with ACS and healthy controls [51,62].

For minimally complex samples such as synthetic peptides or pure compounds, direct infusion mass spectrometry (DIMS) or shotgun lipidomics (for lipid samples) are used. The samples are introduced directly into the MS without prior chromatographic separation. Though suited for high-throughput metabolomics analysis, DIMS often suffers from the matrix effect and ion suppression since all the sample components are infused simultaneously [81,82]. Direct infusion also leads to ion source contamination, which usually takes a long time to recover. Recently introduced chip-based direct-infusion nano-electrospray interfaces were successful in resolving this problem to a great extent [83]. Due to the lack of chromatographic separation, DIMS also cannot separate isomeric compounds. One approach to achieving isomeric separation is incorporating ion mobility spectrometry (IMS) as the separation process before MS [84]. Another drawback of the DIMS system is that multiple ions of the same molecule, such as its molecular ions, adducts, and in-source fragments, can be present in the mass spectrum, thus making interpretation of the data very difficult.

Compared to NMR, MS is the method of choice for global metabolite profiling and identifying unknown compounds within samples. MS can detect various metabolite classes (polar, non-polar, and neutral) depending on the choice of ionization mode (positive/negative). For instance, positive ESI mode works well with medium-sized polar molecules, whereas negative ESI mode is suitable for carbohydrates and organic acids. It is worth mentioning that all clinical studies of lipidomics in ACS mentioned in Table 3 were carried out using the LC/MS platform. Additionally, MS also provides an excellent analytical platform for profiling the molecular composition of lipoprotein complexes. For example, multiple clinical studies [85–87] employed the LC/MS platform to highlight the changes in lipidome composition of HDL (high-density lipoprotein) during ACS pathogenesis.

### Table 3. Main findings from the clinical studies of lipidomics in acute coronary syndrome.

| No | First Author, Year | Sample Size | Sampling Time | Specimen/Technique | Main Findings |
|----|---------------------|-------------|---------------|--------------------|---------------|
| 1  | L. Zhang [88], 2021 | 20 STEMI    | 30 min before PCI; 6, 12, 24, and 72 h after PCI; 1 day before discharge; and 28 days after PCI | Plasma LC/MS | The circulating levels of PGE2, PGD2, and TXA2 were significantly lower at 6 h post-PCI in STEMI patients. The levels of 20-HETE content were significantly higher at 12–72 h post-PCI. |
| 2  | J. Burrello [89], 2020 | 7 STEMI; 9 controls | Pre-PCI, and 24 h post-PCI | Isolated EV Plasma LC/MS | The levels of ceramides, dihydroceramides, and sphingomyelins in extracellular vesicles increased in STEMI compared to matched controls and decreased post-PCI. |
| 3  | P. J. Meikle [90], 2019 | 47 ACS; 83 stable CAD | Before coronary catheterization | Plasma LC/MS | Venous plasma lipid species was better than traditional risk factors in discriminating ACS from stable CAD. |
| 4  | J. H. Lee [87], 2018 | 30 CAD, 10 ACS, 10 with stable CAD without ACS | Not specified | Plasma LC/MS | Two LPC species (16:0 and 18:0) were significantly elevated only in the HDL of the ACS group vs. the stable CAD group, whereas PE species (38:5 and 40:5) were elevated in ACS by >2-fold in both HDL and LDL. |
| Study Reference  | Year | Sample Size | Study Design | Analysis Method | Findings |
|------------------|------|-------------|--------------|----------------|----------|
| Gerl [91], 2018  | 74 ACS, 78 SA, 21 IS, 52 HC | Within the first 24 h of hospital admission | Plasma LC/MS | The levels of LPC and ratios of CE to free cholesterol were decreased in the CVD subjects compared to control subjects. |
| Anroedh [92], 2018 | 581 ACS; 155 MACEs | Prior to coronary angiography or PCI | Plasma LC/MS | The circulating ceramides were associated with MACEs independent of clinical risk factors in CAD patients. |
| Feng [93], 2018 | 40 STEMI | Pre-PCI, 2 h and 24 h post-PCI | Plasma LC/MS | 16 circulating fatty acids were associated with myocardial reperfusion injury. |
| Garcia [94], 2018 | 30 ACS; 30 No CAD | Before hospital discharge | Plasma LC/MS | HDL2 subclass from ACS patients was enriched with oxidized polyunsaturated fatty acids. |
| de Carvalho [95], 2018 | Discovery: 337 Validation: 119 | Pre-angiography and within 24 h post-angiography | Tissue, Plasma LC/MS | 11 ceramides (C14 to C26) and 1 dihydroceramide (C16) were associated with MACEs in patients with AMI. |
| Chatterjee [96], 2017 | 175 symptomatic CAD; 15 HC | During coronary angiography | Platelet LC/MS | Symptomatic CAD patients were characterized by a perturbed platelet lipidome. |
| Zu [97], 2016 | 39 MACE; 39 Non-MACE; 39 controls | During coronary angiography | Plasma LC/MS | The plasma level of 19-HETE is useful for the prognosis of ACS after adjustment for clinical risk factors. |
| Cheng [98], 2015 | 162 STEMI; 151 NSTEMI; 261 stable CAD | Prior to coronary angiography | Plasma LC/MS | Plasma ceramide (d18:1/16:0) was associated with vulnerable plaque and 1-year MACE. |
| Rached [86], 2015 | 16 STEMI; 10 controls | Within 24 h after diagnosis | Plasma LC/MS | The lipidome of HDL particles were markedly altered in STEMI. |
| Sutter [85], 2015 | 23 stable CAD; 22 ACS; 22 HC | Within 12 h of the initial symptom attack | Plasma LC/MS | HDL-plasmalogens levels were inversely associated with both stable and acute CAD. |
| Park [99], 2015 | 140 CAD; 70 HC | After fasting for 12 h | Serum LC/MS | The study showed that multivariate models using multiple lipid species can stratify unstable and stable CAD patients with improved accuracy compared to traditional risk factors. |
| Meikle [100], 2011 | 60 SA; 80 UA; 80 HC | Not specified | Plasma LC/MS | PC containing palmitic acid, DG, SM, and Cer were associated with an increased risk of MI, whereas PE-plasmalogens and PI were associated with a decreased risk. |

Abbreviations: MACE, major adverse cardiac events; STEMI, ST-elevation myocardial infarction; NSTEMI, non-ST-elevation ACS; AMI, acute myocardial infarction; MI, myocardial infarction; SA, stable angina pectoris; UA, unstable angina pectoris; IS, ischemic stroke; PCI, percutaneous coronary intervention; ACS, acute coronary syndromes; CAD, coronary artery disease; HC, healthy control; LC/MS, liquid chromatography/mass spectrometry; LPC, lysophosphatidylcholine; PE, phosphatidylethanolamine; CE, cholesteryl ester; PC, phosphatidylcholine; DG, diacylglycerol; SM, sphingomyelin; Cer, ceramide; PI, phosphatidylinositol; HDL, high-density lipoprotein.

NMR and MS primarily use two strategies to conduct metabolomics and lipidomics studies, i.e., untargeted, and targeted approaches [101,102]. The untargeted approach, also called open profiling, aims to analyze the most comprehensive set of metabolites from biological samples and is non-selective and non-discriminative towards metabolites screening. It is mainly used for the discovery of novel biomarkers or to generate a specific hypothesis. Usually, no pre-existing knowledge about the metabolic profile of the sample is required for this type of analysis. The untargeted approach usually compares metabolite levels between different groups (e.g., healthy control vs. disease) under similar conditions [101,103]. A schematic showing the different steps in untargeted metabolomics is shown in Figure 3.

On the other hand, the targeted approach or closed profiling focuses on a limited number of known metabolites, such as 10–20 lipids, and seeks to quantify them. A hypothesis is being tested in this approach, and often the sample preparation and analytical
techniques are more sophisticated. In the case of ACS, the most used approach so far has been untargeted metabolomics to differentiate the ACS profile from healthy controls. However, most ACS studies complement their untargeted analysis by performing a targeted analysis of those compounds that exhibited a significant difference between various groups for further quantitation and identity confirmation using the suitable internal standards. In a study to understand the underlying mechanisms associated with CAD progression, Zhang and colleagues initially performed an untargeted analysis of metabolites in plasma of 2324 patients who underwent coronary angiography [61]. They identified a total of 36 differential metabolites across different CAD types, including N-acetylneuraminic acid (a ligand for many hormones and lectins), whose levels were elevated in plasma during CAD progression. Subsequently, a targeted quantification was performed using isotope-labeled N-acetylneuraminic acid to confirm its vital role in CAD progression.

Figure 3. A standard workflow for untargeted metabolomics. After sample collection and extraction, the samples are analyzed by NMR spectroscopy or mass spectrometry. The raw data are then analyzed using appropriate software followed by statistical analysis to identify metabolites of interest or potential candidate biomarkers. Adapted from “Untargeted Metabolomics for Discovery of Disease Biomarkers”, by BioRender.com (2021). Retrieved from https://app.biorender.com/biorender-templates on August 11, 2021.

3. Pre-Analytical Considerations in Metabolomics Studies

The analytical quality of both untargeted and targeted metabolomics approaches is mainly dependent on the various pre-analytical steps involved in sample handling, sample collection, centrifugation, aliquoting, transportation, freezing, and storage [104]. Even minor discrepancies in any of these pre-analytical factors can greatly influence metabolomic assessments. Discrepancies may include, but are not limited to, choice of anticoagulants or preservatives added to the sample specimen, time delay, and storage temperature during blood preprocessing [105], diet and time of sampling in urinalysis [106,107], metabolite degradation or aggregation induced by air oxidation and light exposure [108,109], number of freeze–thaw cycles before aliquoting biological fluids [110], and time delay in transportation and sample storage [111]. Therefore, knowledge about different
sample collection procedures and preparation protocols is essential to ensure reliable and reproducible results in a metabolomics study. A variety of biological fluids, including plasma, serum, urine, saliva, cerebrospinal fluid, bronchoalveolar lavage fluid, and cell and tissue extracts, are used in metabolomics studies depending upon the biological question under study and the pathophysiological nature of the disease process [112].

3.1. Serum vs. Plasma

Notably, most of the metabolomics studies in ACS have been performed using either plasma or serum as the sample source [33,34,69–71]. The general theme of most of these plasma- or serum-based studies is to find biomarkers associated with ACS and its subtypes, compared to healthy controls. Though both plasma and serum matrices provide an excellent functional readout of an organism’s metabolic activity, the question regarding which is more suitable is still highly debatable. Both plasma and serum are derived from the liquid portion of the blood. The serum is extracted after the blood has clotted, and it is obtained by centrifugation after clotting. For plasma, the blood is not coagulated and is obtained by adding an appropriate anticoagulant (heparin, citrate, or Ethylenediaminetetraacetic acid (EDTA)) to the whole blood, followed by centrifugation and aliquoting [112].

Typically, plasma makes up to 55% of the total volume of blood, and it contains dissolved proteins, clotting factors, salts, lipids, and other suspended materials in water. The serum is similar to plasma, except it lacks fibrinogen (clotting factor). Contrary to serum, plasma has the advantage that it can be immediately placed on ice, thereby avoiding the possible loss of labile metabolites due to enzymatic conversion or other degradation processes during clot formation at room temperature [112]. Several studies have tested the role of pre-analytical factors on the metabolic composition of serum and plasma [113–115]. Teahan et al. showed that variations in certain pre-analytical factors such as clotting time and temperature, the presence/absence of anticoagulant, and storage conditions including repeated freeze−thaw cycles, can introduce bias into metabolic data [115]. Additionally, recent data suggest that it is not advisable to combine serum samples exposed to different clotting procedures (e.g., thrombin vs. silicate-enhanced) or different clotting times into a single sample set for biomarker analysis, as coagulation and associated processes can alter metabolite concentrations [116]. This limitation can be highly challenging, particularly in multicentric clinical studies involving humans, where samples are collected from different hospital sites and transported to a centralized facility for further processing and analysis.

Besides assessing various pre-analytical factors, numerous studies have also investigated the difference in metabolic profile generated by serum and plasma samples [116–118]. In a study comparing serum and plasma metabolic profiles [116], it was found out that 46% out of the 216 identified metabolites were significantly different between the two, and except for three (methionine, C2:0-, and C3:0-carnitine), the levels of all other metabolites were higher in serum. In line with this, in another metabolomics study comprising 377 individuals [118], it was reported that although the reproducibility was slightly better in plasma than serum, the concentrations of analytes were generally higher in serum, with an average relative difference of 11.7%. In another study involving 29 small-cell lung cancer patients [117], it was shown that neither fluid is superior to the other, and the two metabolomes were markedly similar in terms of reproducibility, specificity, and metabolic coverage. These studies suggest that even though both biofluids are comparable in terms of analytical throughput, serum may provide better sensitivity than plasma.

3.2. Polar vs. Non-Polar Metabolites

Due to the enormous diversity in the physicochemical properties of metabolites, especially polar (including amino acids, nucleic acids, and sugars) and non-polar (including fatty acids and other lipids) metabolite classes, no ideal extraction protocol exists that can extract both polar and non-polar metabolites from the same sample. Accordingly, a comprehensive analysis of the entire metabolome in a biological sample requires multiple extraction strategies to cover different metabolite classes. Hence, multiple aliquots of the
same sample with specific extraction procedures, such as using a modified Bligh–Dyer protocol for extracting polar metabolites [119], are required to obtain extensive coverage of the entire cellular metabolome, which in turn facilitates the need of a larger sample amount. In studies where sample amount is a limiting factor, this approach might not be feasible. Of late, efforts have been made to develop extraction protocols that can extract both polar and non-polar analytes from the same sample [120]. Selecting the proper extraction protocol is critical in all metabolomics studies as the metabolite extraction step directly affects all downstream steps of metabolomics data analysis.

4. Extraction Procedures for Metabolomics

The reliability and accuracy of metabolic data also depend upon the sample preparation strategy. Ideally, the sample processing should be fast, easy, minimal, and tailored to the analytical scheme. Common approaches include homogenization, dialysis, fractionation, extraction, distillation, centrifugation, and concentration. Critically, these approaches should be compatible with the nature of the biological matrix, chemical/physical properties of the analyte of interest, and the final detection technology. For example, in a typical LC/MS platform, sample preparation is usually the most time-consuming and error-prone step of the chromatographic assay. The inherent ‘matrix effect’ on different samples such as plasma, serum, urine, and tissue lysate, presents significant analytical challenges during LC/MS analysis [121]. Matrix effects are caused by co-eluting endogenous components and preservative agents in the same matrix [122], which often lead to material buildup on the analytical column and ion source in LC and MS, respectively. This can cause ion suppression or enhancement, drift in chromatographic response, increased or reduced analytical signal, reduced column life, and frequent MS cleaning [123]. Cumulatively, these issues may compromise analytical accuracy and increase the total analysis time, in addition to significantly affecting the cost of MS analysis [121]. Therefore, thoughtful selection and optimizing sample preparation procedures are essential to minimize variability and improve analytical performance in metabolomics studies.

The three most popular sample preparation techniques in metabolomics analysis include protein precipitation (PPT), liquid–liquid extraction (LLE), and solid-phase extraction (SPE) [124,125]. PPT generally begins with adding organic solvents such as methanol, acetonitrile, or a combination thereof to the sample, followed by agitation and centrifugation [126]. Its ‘nonselective’ nature makes it well-suited for global metabolomics analysis. Adding an ice-cold organic solvent is beneficial as it may improve the efficiency of protein removal and prevent metabolite degradation as the sample warms up from its stored frozen state [127]. While agitation increases the protein accumulation rate, centrifugation helps separate the supernatant holding analytes from the protein pellets [128]. Though this method provides high metabolite coverage, PPT is often time-consuming, particularly while dealing with hundreds of samples manually, as in large-scale epidemiological studies. Robotic systems capable of performing automated protein precipitation such as membrane-based protein precipitation filter plates have been developed recently to address this issue [124,129,130].

Another classic method used for the qualitative/quantitative identification of metabolites is biphasic LLE. LLE utilizes two immiscible liquids for the extraction of analytes. In LLE, the analyte is differentially distributed between the aqueous matrix and water-immiscible organic solvent [124]. Traditionally, LLE is most preferred for comprehensive lipid analysis [131] (i.e., global lipidomics). Apart from enriching analytes of interest (here, lipids) and improving signal-to-noise ratios, any extraction protocol used in lipidomics should also remove any non-lipid compounds. Being insoluble in water, lipids are extracted from a biological matrix, such as blood plasma or tissue, using organic solvents. The most widely used method for isolating lipids from biological samples was devised by Folch et al. more than 60 years ago using 2:1 chloroform/methanol (v/v) as the solvent mixture [132]. In this two-phase LLE procedure, most lipids are dispersed into the lower chloroform phase, and are clearly separated from the upper methanolic phase holding
non-lipid substances. To enhance recovery, Bligh and Dyer later revised Folch’s method by combining chloroform, methanol, and water [133]. The disadvantage of these two methods is that the recovery of the lower lipid fraction is often cumbersome, resulting in contamination of the isolate and blockage of the analytical column. In 2008, Matyash and co-workers [134] showed that an extraction procedure using methyl-tert-butyl ether (MTBE) could solve this problem as the lipid-containing organic layer settles at the top during phase separation, thereby enabling a much cleaner lipid extraction. More recently, a chloroform-free one-phase lipid extraction protocol based on a mixture of butanol and methanol (BUME) was described by Löfgren and coworkers [135]. This novel approach is rapid and shows similar extraction efficiency compared to the Folch and the Bligh and Dyer procedures.

In SPE, samples are loaded onto a solid sorbent held primarily on a cartridge device (SPE cartridge). The analyte of interest present in samples is selectively retained by the sorbent material [136]. The retained analyte is then eluted with a suitable solvent [137]. Different SPE types include reverse, normal, anion, cation, and mixed-mode sorbents [124]. Due to its highly selective nature, SPE is not recommended for global metabolomics profiling [138]. It is mainly used to concentrate analytes present at low levels and purify analytes from matrix interferences. There exist contradictory reports regarding the analytical reproducibility using the SPE procedure. In a study comparing PPT and SPE procedures, SPE showed improved repeatability compared to PPT for human plasma metabolic profiling [139]. However, in another similar study evaluating different human plasma preparation protocols, the metabolome coverage and repeatability of the SPE procedure are reported to be lower than PPT protocols [126]. Once an extraction procedure is selected, caution should be taken to ensure that the same procedure is repeated consistently throughout the entire study. For large-scale cohort studies, maintaining this consistency presents a real challenge. Inconsistency in extraction procedures is a source of variation in the dataset and may compromise the robustness and accuracy of the results.

5. Data Processing in Metabolomics

As with the genomics platform, metabolomics studies generate a large amount of multidimensional, non-linear, and non-normal data [140]. Due to its sheer complexity, the metabolomics data-analytical approach relies heavily on advanced computational approaches. Many instrument-specific and open-source software solutions are available, such as MetaboAnalyst [141] and XCMS [142], that can perform the critically important steps in metabolomics data analysis, such as run alignment, peak picking, data preprocessing (e.g., deconvolution, scaling, and normalization), annotation, and compound identification. However, selecting a suitable approach or software is often confusing and depends upon the separation technique, detection mode, and output file format (e.g., mzXML, mzML, NetCDF) generated by each instrument. Additionally, there remains little accord or harmonization among different software in handling various cross-vendor file formats. Recently, much effort has been made to address this issue, and new software platforms such as Progenesis QI [143] have been developed, that are compatible with a wide variety of instruments and can perform platform-independent metabolomics data analysis.

5.1. Handling Unwanted Variances in Metabolomics Data

Analytical and biological variability issues are critical in human metabolomics studies. If these unwanted variations are not accounted for, it may affect the statistical power of detecting metabolites that are characteristic of, for example, a disease state such as ACS. Laboratory procedures or technological platforms primarily introduce analytical variability. Typical examples include matrix effects, temperature changes, solvent pH changes, sample degradation over time in extended sample runs spanning weeks or months, fluctuation in instrumental sensitivity, and personal errors [144]. One way to combat these analytical errors is by employing suitable sample normalization strategies, such as data-
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 driven normalizations, internal standards-based normalization, or quality control (QC-based normalization [145]. These strategies help to standardize metabolite abundances before statistical analyses and improve the quality of metabolomics data. Of these, QC-based normalization approaches are gaining more popularity of late [144,146]. Several QC-based normalization algorithms have been developed in recent times, including batch-ratio [147], LOESS [148], and SERFF [145]. The QC samples are injected at regular intervals along with the study samples in every batch. Ideally, the QC sample should have the same matrix composition as that of study samples and is usually obtained by pooling multiple aliquots. After QC-based batch correction, those metabolites which exhibited poor repeatability across QC samples were often removed based on specific cut-off criteria to assure an expected level of data quality. For example, after batch correction, only those metabolites satisfying the cut-off criteria were often retained for subsequent data analysis [149], such as (1) metabolites present in >50% of QC samples, and (2) metabolites with a coefficient of variation for QC samples <20%.

Though the above-mentioned approaches can largely eliminate unwanted analytical variations from the dataset, the inherent biological variability induced by factors such as diet, ethnicity, physical activity, circadian rhythm, and medications still poses a significant challenge in interpreting the real physiological variations due to disease status or interventions [150]. To overcome these challenges, several strategies have been proposed, including cross-validation of results using an appropriate proportion of test and validation sets (e.g., 30%, the ‘test set’), validation of identified biomarkers using large cohorts of multi-centric samples (validation set), reporting results in relative amounts rather than absolute amounts of metabolites, and finally, validation of candidate biomarkers based on comparison to isotopically labeled internal standards. Current practice involves using different software for different tasks in the metabolomics workflow, such as one for peak picking, another one for normalization, and another one for metabolite annotations. This is a very tedious process and a time-consuming one. As we move forward, particular emphasis should be given to developing software that can automate this process from aligning runs to peak picking to batch correction to metabolite annotations, which may become “current practices” in the immediate future.

6. Metabolomics in ACS

Recently, there has been a growing appreciation for metabolomics as a promising approach for investigating cardiovascular diseases, including CAD to allow for better mechanistic understanding and biomarker discovery. The studies in the 2000s by Sabatine and colleagues initially demonstrated the application of a blood-based metabolomics platform to identify markers associated with myocardial injury. In a study comprising 36 subjects who underwent exercise stress testing (inducible ischemia), Sabatine et al. showed that myocardial ischemia is characterized by significant changes in the circulating levels of multiple metabolites, including lactic acid (final product of glycolysis) [151]. The pathway analysis identified a key role for the TCA cycle (regulator of oxidative phosphorylation) during myocardial ischemia. Moreover, they identified a panel of six metabolites, including citric acid, which could accurately stratify patients with myocardial ischemia from control subjects. Following this study, in a human model of planned myocardial infarction, via alcohol septal ablation, the same group identified metabolic changes as early as 10 minutes after myocardial injury [152]. They reported that perturbations in pyrimidine metabolism, the TCA cycle, and the pentose phosphate pathway were associated with myocardial injury. Importantly, these perturbations were observed in both coronary sinus and peripheral blood and were further validated in an independent clinical cohort.

A comprehensive overview of the clinical studies of metabolomics in ACS is provided in Table 2. Only those studies which explicitly discuss any of the ACS subtypes were included in the table. As evident from Table 2, blood-derived plasma and serum from veins are the most preferred biological fluids, and LC/MS is the favored technique being used in these studies. However, the sampling time varies greatly between these
studies. First-morning collection, collection after overnight fasting, spot sampling at the time of admission (non-fasting), and timed collection at different sampling time points are the various sampling modes employed in these studies. The ideal sampling time is when the rate of metabolic flow, or flux, is constant, i.e., a steady metabolic state. However, ACS always involves ‘metabolic shifts’, and thus there is no steady metabolic state available in ACS events. Under these circumstances, time-series analyses could provide a better insight into the molecules and pathways with clinical relevance.

The initial clinical studies of metabolomics in ACS made use of GC/MS to identify biomarkers for the early diagnosis of ACS. Utilizing GC/MS, Vallejo et al. and Laborde et al. elucidated the metabolic differences in plasma of non-ST-elevation ACS patients relative to healthy subjects [68,71]. These studies collectively identified tricarboxylic acid (TCA) cycle intermediates along with 4-hydroxyproline, tryptophan, 3-OH-butyric acid, and 2-OH-butyric acid as key players in ACS pathophysiology. Subsequent studies employed NMR, LC/MS, and CE/MS approaches, shifting focus from the broad spectrum of ACS towards various ACS subtypes. Using CE/MS and hydrophilic interaction chromatography/MS-targeted analysis, Naz et al. found increased acylcarnitines (associated with defective mitochondrial β-oxidation) and amino acids (involved in myocardial energy metabolism) levels in STEMI patients compared to NSTEMI patients [67]. Another study used a combination of different metabolomics approaches, including GC/MS and H-NMR, and confirmed the presence of elevated hydrogen sulfide (an endogenous gasotransmitter) levels in STEMI patients compared to UA patients [63]. In a large study comprising 2324 patients from 4 independent centers [64], Fan et al. evaluated the diagnostic value of plasma metabolomics to characterize different types of CAD. Based on CAD severity, patients were divided into five groups, of those with the normal coronary artery, nonobstructive coronary atherosclerosis, stable angina, UA, and AMI. They found 89 differential metabolites across different CAD types. Additionally, they identified glycerophospholipid metabolism, amino acids, acylcarnitines, TCA cycle, and bile acid biosynthesis as the main metabolic pathways associated with CAD progression. Importantly, these findings were replicated in a validation cohort.

Metabolomics has also been used to explore biomarkers predictive of adverse cardiovascular events following ACS. For instance, Du et al. performed LC/MS analysis of 26 amino acids in a cohort of 138 STEMI patients with acute heart failure to find metabolites predictive of adverse cardiovascular events [58]. They found that elevated plasma branched-chain amino acids (BCAA) levels on admission are associated with adverse cardiovascular events. In another study comprising 978 patients [54], Vignoli et al. used NMR-based metabolomics to identify prognostic markers of two-year mortality after AMI. They showed that elevated levels of amino acids including mannose, formate, acetone, proline, creatinine, acetate, and 3-hydroxybutyrate were associated with mortality following AMI. Another study used an untargeted LC/MS approach and showed that six metabolic pathways, namely urea cycle, tyrosine, lysine, tryptophan, aspartate/asparagine, and carnitine shuttle, are associated with mortality in patients with CAD [47]. More recently, Chorell et al. showed that lysophospholipids (involved in inflammation and arteriosclerosis) are associated with future cardiovascular risk in STEMI and NSTEMI patients [44]. They reported that while STEMI is characterized by a higher ratio of lysophosphatidylcholine to lysophosphatidylethanolamine, NSTEMI is characterized by a lower ratio of these two lipids [44].

In recent times, lipid molecules and their associated pathways gained particular interest in the setting of ACS. In 2018, Wang et al. demonstrated that in addition to TCA cycle intermediates and amino acid metabolism, other lipid-associated pathways, including fatty acid metabolism and fatty acid β-oxidation, also play important roles in ACS [56]. During the same time, Goulart et al. showed that the most perturbed metabolites associated with STEMI were primarily lipid species, including phosphatidylcholines, lysophosphatidylcholines, and sphingomyelins [55]. These results underscore the need for comprehensive lipid profiling to provide insight into ACS pathogenesis.
One of the most consistent findings in these clinical studies has been the link between carnitine (short-chain and long-chain) and lysophosphatidylcholine (LPC) species with ACS. Carnitines play a critical role in transporting long-chain fatty acids from the cytoplasm into the mitochondria, where they undergo β-oxidation to produce energy. Accumulating evidence suggests that elevated levels of carnitines reflect impaired β-oxidation and mitochondrial dysfunction [153] and are associated with a wide variety of disorders, including type 2 diabetes [154], and cardiovascular diseases [155]. Since long-chain fatty acylcarnitines are produced from fatty acid metabolism and are primarily synthesized in the mitochondria, their levels indicate mitochondrial fatty acid oxidation [156]. On the other hand, LPC is a group of proinflammatory lipids, which is primarily derived from phosphatidylcholine (PC) by the enzymatic action of phospholipase A2 (PLA2). LPC has been linked to the pathogenesis of atherosclerosis and the progression of various diseases [157], including cardiovascular diseases, renal failure [158], ovarian cancer [159], and diabetes [160]. Among other properties, LPC also activates several signaling pathways, such as oxidative stress and inflammatory responses, contributing to endothelial cell injury in atherosclerosis and cardiovascular disorders [161].

7. Lipidomics in ACS

Apart from their primary role as the structural components of cells, lipids exert indispensable functionalities as cell signaling molecules and energy sources. Evidence from genomics studies and large randomized controlled trials has established the link between the dysregulated lipid metabolism and CAD progression, including ACS. The results from clinical trials of lipid-modifying therapy demonstrated that lowering the levels of serum lipids (especially cholesterol) reduces the risk of cardiovascular events [162]. Given the crucial role of lipids in regulating health and disease states, elucidating the lipid composition at the molecular and system level is essential to characterize the molecular basis of ACS. Therefore, there is a growing interest in lipidomics as a promising approach to reveal lipid alterations in ACS progression and to find new biomarkers for early ACS diagnosis. Table 3 summarizes the applications of lipidomics on ACS in clinical settings. From Table 3, it is evident that MS has been the dominating technique for lipid profiling, and blood-derived plasma is the most preferred biospecimen.

Traditional clinical lipid biomarkers for the development and progression of CAD, including elevated serum low-density lipoprotein (LDL), decreased high-density lipoprotein (HDL), or increased triglycerides levels, often fail to correctly distinguish ACS from stable coronary artery disease. Using plasma lipid profiling on 220 individuals, Meikle and colleagues showed that multivariate models incorporating both lipids and conventional risk factors could stratify unstable CAD from stable CAD patients with better accuracy than models with conventional risk factors alone [100]. The plasma levels of many lipids, including alkylphosphatidylcholine and phosphatidylcholine plasmalogen (these species are susceptible to oxidative stress), displayed a significant association with disease severity, suggesting their part in the onset and progression of ACS. In a report to characterize lipid species within lipoprotein particles, Meikle also reported that the levels of phospholipids, including lysophospholipids and plasmalogen, were significantly lower within the HDL of the ACS group relative to the CAD group [90]. In line with this finding, Sutter et al. and Rached et al. showed the contribution of alterations in the HDL lipidome to the disease severity of ACS [85,86]. Similarly, looking at 365 lipids, Lee et al. demonstrated that the levels of saturated lysophosphatidylcholine (LPC) species (16:0 and 18:0) were increased only in the HDL fraction of the ACS group, indicating an intermediate link between LPC species and progression of ACS [87]. In another study using a targeted lipidomics approach, Garcia et al. showed that the HDL2 subclass of ACS patients is enriched with oxidized fatty acids compared to non-ACS subjects, which may modulate platelet-dependent thrombotic risk [94]. Together, these studies demonstrate the ability of whole-plasma and lipoprotein-specific lipidomics for the early detection of ACS and discriminating stable CAD from ACS.
Recently, several studies have looked at the association between molecular lipid species and clinical outcomes in patients with ACS. In a study comprising 581 patients with ACS or stable CAD, Cheng et al. investigated the association of plasma lipids with 1-year clinical outcome [98]. They showed that plasma concentration of ceramides (involved in inflammation, membrane integrity, and apoptosis), particularly Cer(d18:1/16:0), is strongly associated with 1-year major adverse cardiac events (MACE) and plaque vulnerability, independent of statin usage and LDL levels. The prognostic value of these high-risk circulating ceramide species was further probed in the prospective ATHEROREMO cohort of 581 patients with stable angina pectoris or ACS and a median follow-up of 4.7 years [92]. Multivariable analyses showed that the circulating levels of Cer(d18:1/16:0), Cer(d18:1/20:0), Cer(d18:1/24:1), and their ratios were associated with adverse cardiac outcomes independent of the established clinical risk factors. A recent lipidomics analysis by Carvalho et al. employing paired tissue–plasma samples in human and animal models took these results a step further [95]. They showed that arterial and myocardial tissue ceramide levels also correlate with MACE in patients with AMI. Collectively, these data suggest a predictive role of plasma ceramide species in patients with ACS.

8. Metabolomics of Ischemia/Reperfusion Injury

Timely myocardial reperfusion strategies employing fibrinolytic therapy or percutaneous coronary intervention (PCI) are the treatment of choice for acute STEMI patients. Besides salvaging viable cardiomyocytes from ischemic death, reperfusion profoundly limits the infarct size following a prolonged ischemic insult and improves the clinical outcome. However, pre-clinical and clinical data show that this sudden reintroduction of oxygen and nutrients during reperfusion by itself induces cardiomyocyte death, a phenomenon termed myocardial reperfusion injury [13,163]. Clinically, the largest effect of ischemia/reperfusion (I/R) injury is during PCI of STEMI patients. In STEMI patients, once the coronary artery is opened by PCI (Figure 4), it allows for reperfusion of the ischemic myocardium. Despite rapid and successful reperfusion, the mortality rate after an AMI is nearly 10% [164].

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**Figure 4.** Percutaneous coronary intervention (PCI): PCI widens the blocked or narrow coronary arteries, thereby allowing reperfusion of the ischemic myocardium. Adapted from “Percutaneous Coronary Intervention”, by BioRender.com (2021). Retrieved from https://app.biorender.com/biorender-templates on August 11, 2021.
Low blood flow during acute ischemia switches cell metabolism to anaerobic metabolism, resulting in lactate accumulation and lowering of intracellular pH [165]. Additionally, there is a decrease in cellular ATP, which results in Ca²⁺ overload [165]. These processes disturb cell volume regulatory mechanisms, leading to disrupted cellular structure and cell lysis. Thus, it is essential to restore tissue oxygen supply. However, reperfusion leads to a sudden increase in the amount of oxygen available that causes an intense burst of mitochondrial reactive oxygen species (ROS), resulting in cellular dysfunction by modifying intracellular molecules [165]. Reperfusion also restores the physiological pH, which releases the inhibitory effect on mitochondrial permeability transition pores’ (mPTP) opening. Moreover, reperfusion results in intracellular Ca²⁺ overload due to the dysfunction of the sarcoplasmic reticulum. Reperfusion also initiates endoplasmic reticulum stress, a pro-inflammatory response, and pro-thrombogenic pathway activation in ischemic tissues [163,165]. The excess Ca²⁺ and enhanced ROS production trigger mPTP opening, resulting in ATP depletion and, ultimately, cellular death. A graphic indicating the main proponents of myocardial I/R injury is shown in Figure 5. In brief, reperfusion is associated with an additional injury that extends the ischemic damage. As a result of this puzzling problem between ischemic injury vs. reperfusion injury, there is significant interest in finding ways to protect against and treat the latter.

Figure 5. Myocardial ischemia/reperfusion (I/R) injury. Schematic showing the main events in myocardial I/R injury. Abbreviations: Ca²⁺, Calcium ion; ROS, reactive oxygen species; mPTP, mitochondrial permeability transition pore.

The ROS generation during reperfusion was previously thought of as a non-specific response to reoxygenation of ischemic tissue. However, a recent comparative in vivo metabolomic analysis in a mouse model of I/R injury has revealed that the TCA cycle intermediate, succinate accumulates during ischemia and the oxidation of succinate during reperfusion drives mitochondrial ROS accumulation and reperfusion injury [166]. A later study used a targeted LC/MS approach to analyze plasma metabolites in 115 STEMI patients undergoing PCI and revealed that myocardial succinate accumulation is an early marker of human I/R injury [60]. They reported an association between myocardial content of succinate and the magnitude of ischemic injury in STEMI patients.
Recent metabolomics studies have also implicated the role of BCAAs in the development of I/R injury. In a metabolomics study utilizing a KO mouse model (2C-type ser/thr protein phosphatase (PP2Cm) deficient), Li et al. showed that impaired BCAA catabolism suppresses glucose uptake and worsens the I/R injury [167]. In line with this finding, our previous study in a clinical setting of human I/R injury identified BCAA (leucine, isoleucine, and valine) biosynthesis as one of the most perturbed metabolic pathways during early reperfusion [52]. Our study showed that lipids and their derivatives formed the bulk of the altered metabolome in the setting of reperfusion injury in STEMI patients [52]. We also identified a panel of three lipid molecules, namely pentadecanoic acid, 18:2 carnitine, and 18:2 lysophosphatidylcholine, that can determine the extent of I/R injury in STEMI patients after primary PCI.

A few studies have also explored the time-course changes in the circulating metabolic profile following reperfusion. Based on a time-series analysis before and after PCI comprising 40 STEMI patients, Feng et al. showed that the fatty acid content in the circulating blood gradually decreases with an increase in reperfusion time [93]. In another study monitoring the changes in eicosanoid profile before and after PCI in 20 STEMI patients, Zhang et al. found that the levels of some bioactive eicosanoids, including PGE2, PGD2, TXA2, and 20-HETE, were significantly decreased after PCI. Interestingly, these molecules are pro-inflammatory and are associated with platelet aggregation [168].

There are currently no therapeutic options for I/R injury in patients presenting with STEMI who undergo revascularization. Given the complex metabolic changes within the myocardium and hence within the circulation during I/R injury, it is necessary to identify the metabolic and lipidomic pathway(s) that impact clinical outcomes. There are many limitations to the current studies in the setting of I/R injury, including focusing on a single metabolite (e.g., succinate), a specific metabolite class (e.g., fatty acids or eicosanoids), a single pathway (e.g., BCAA metabolism), or having small sample sizes. Animal studies and planned strategies (e.g., inducible ischemia) are helpful to understand pathological mechanisms. Nevertheless, more clinical investigations incorporating time course analyses in large cohorts are needed to better understand the dynamic changes and metabolic pathways involved in human I/R injury.

9. Translational Metabolomics and Future Directions

Despite showing substantial potential for biomarker discovery and a more detailed understanding of pathogeneses of ACS, few findings from metabolomics studies have been translated into disease diagnostics and risk prediction. One main reason is the validity and reliability of new markers/clusters of metabolites [169]. The main challenge in biomarker validation is the difficulty in measuring subtle differences in metabolite fluctuations associated with different disease status or interventions. Additionally, as evident from Tables 2 and 3, barring a few, most of the clinical metabolomics studies on ACS are small-scale preliminary-type studies performed with a limited sample size ($n < 100$), which lack the required statistical robustness and validity. We believe the only way to overcome this is to perform large clinical studies to allow for successful translation of metabolomics to clinics. In addition, as discussed in Section 5.1, the inter-individual variation induced by diet, genetic, and environmental exposures, coupled with technical variability, also diminishes metabolomics studies’ power to detect actual physiological variations associated with different ACS subtypes and, eventually, to provide clinical biomarkers [170]. One way to overcome this is by conducting follow-up metabolomics experiments using a large validation cohort consisting of diverse patient groups (multi-centric) with suitable control cohorts. However, large-scale multi-cohort studies are usually laborious, expensive, and time-consuming. Once validated appropriately, there are increased prospects of translating these biomarkers towards clinics and diagnostic centers. Another potential challenge in translational metabolomics is that most of the metabolomics studies in ACS published thus far were focused on generating data and interpreting them without delving into the mechanistic link underlying the association between the
identified metabolites and ACS pathogeneses [171]. Finding the relation between candidate biomarkers and their biological role is the next important step after biomarker discovery. This can be carried out by employing suitable in vitro and animal model experiments. Importantly, in the future, the metabolomics community should work in tandem with other omics communities such as genomics and transcriptomics to gain more insights on cellular processes represented by a candidate marker [172]. This will bring us closer to a mechanistic understanding of various physiological and pathophysiological conditions associated with various disease states, including ACS.

10. Conclusions
Over the past decade, metabolomics has become a powerful investigative tool to elucidate the underlying metabolic mechanisms of cardiovascular diseases. Most of the metabolomics studies in ACS have focused on biomarker identification to differentiate ACS from healthy controls, to differentiate ACS subtypes (UA, NSTEMI, and STEMI), and to find predictive molecules of mortality or adverse events following an ACS incident. Several studies also performed pathway analysis for finding the biological pathways that contribute to disease pathogenesis. Many of these studies have already shown substantial potential for discovery and understanding. However, as discussed above, attempts to translate these study results into clinical practice have resulted in contradictory results. Considering metabolomics is still early in its scientific evolution, the future is promising with the ongoing technological advances in the field. More importantly, more clinical studies aiming at a system-wide understanding of ACS pathogenesis rather than risk prediction models are the need of the hour.

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Abbreviations
ACS: acute coronary syndrome; AMI, acute myocardial infarction; BCAA, branched-chain amino acids; CAD, coronary artery disease; CE/MS, capillary electrophoresis and mass spectrometry; CVD, cardiovascular disease; GC/MS, gas chromatography and mass spectrometry; I/R, ischemia/reperfusion; LC/MS, liquid chromatography and mass spectrometry; LDL, low-density lipoprotein; HDL, high-density lipoprotein; MACE, major adverse cardiac events; MS, mass spectrometry; NMR, nuclear magnetic resonance; NSTEMI, non-ST-segment elevation myocardial infarction; PCI, percutaneous coronary intervention; ROS, reactive oxygen species; STEMI, ST-segment elevation myocardial infarction; UA, unstable angina.

References
1. Sanchis-Gomar, F.; Perez-Quilis, C.; Leischik, R.; Lucia, A. Epidemiology of coronary heart disease and acute coronary syndrome. Ann. Transl. Med. 2016, 4, 256–256, doi:10.21037/atm.2016.06.33.
2. Virani, S. S.; Alonso, A.; Benjamin, E. J.; Bittencourt, M. S.; Callaway, C. W.; Carson, A. P.; Chamberlain, A. M.; Chang, A. R.; Cheng, S.; Delling, F. N.; et al. Heart disease and stroke statistics-2020 update: A report from the American Heart Association. Circulation 2020, 141, e139–e596, doi:10.1161/cir.0000000000000757.
3. Amsterdam, E. A.; Wenger, N. K.; Brindis, R. G.; Casey, D. E., Jr.; Ganiats, T. G.; Holmes, D. R., Jr.; Jaffe, A. S.; Jneid, H.; Kelly, R. F.; Kontos, M. C.; et al. 2014 AHA/ACC guideline for the management of patients with non-ST-elevation acute coronary syndromes: A report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. J. Am. Coll. Cardiol. 2014, 64, e139–e228, doi:10.1016/j.jacc.2014.09.017.
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4. Menzin, J.; Wygant, G.; Hauch, O.; Jackel, J.; Friedman, M. One-year costs of ischemic heart disease among patients with acute coronary syndromes: Findings from a multi-employer claims database. Curr. Med. Res. Opin. 2008, 24, 461–468, doi:10.1185/030079908x261096.

5. Anderson, J.L.; Morrow, D.A. Acute myocardial infarction. N. Engl. J. Med. 2017, 376, 2053–2064, doi:10.1056/NEJMra1606915.

6. Moore, A.; Goerne, H.; Rajah, P.; Tanabe, Y.; Saboo, S.; Abbara, S. Acute myocardial infarct. Radiol. Clin. N. Am. 2019, 57, 45–55, doi:10.1016/j.rcl.2018.08.006.

7. Makk, N.; Brennan, T.M.; Girotra, S. Acute coronary syndrome. J. Intensive Care Med. 2015, 30, 186–200, doi:10.1177/0885066613503294.

8. Mahajan, V.S.; Jarolim, P. How to interpret elevated cardiac troponin levels. Circulation 2011, 124, 2350–2354, doi:10.1161/CIRCULATIONAHA.111.23697.

9. Taegtmeyer, H. A Primer on Carbohydrate Metabolism in the Heart.; Springer: New York, NY, USA, 2014; Volume 11.

10. Drosatos, K.; Goldberg, I.J. Lipoproteins: A source of cardiac lipids. In Cardiac Energy Metabolism in Health and Disease; Springer: New York, NY, USA, 2014.

11. Burnett, J.R. Lipids, lipoproteins, atherosclerosis and cardiovascular disease. Clin. Biochem. Rev. 2004, 25, 2.

12. Rosano, G.M.; Fini, M.; Caminiti, G.; Barbaro, G. Cardiac metabolism in myocardial ischemia. Curr. Pharm. Des. 2008, 14, 2551–2562, doi:10.2174/138161208786071317.

13. Yellon, D.M.; Hausenloy, D.J. Myocardial reperfusion injury. N. Engl. J. Med. 2007, 357, 1121–1135, doi:10.1056/NEJMra071667.

14. Riehle, C.; Abel, E.D. Insulin signaling and heart failure. Curr. Pharm. Des. 2007, 13, 2374–2376, doi:10.1016/j.cpcpe.2007.04.003.

15. Griﬃn, J.L.; Atherton, H.; Shockey, J.; Atzori, L. Metabolomics as a tool for cardiac research. Nat. Rev. Cardiol. 2011, 8, 630–643, doi:10.1038/nrcardio.2011.138.

16. Dunn, W.B.; Goodacre, R.; Neyses, L.; Mamas, M. Integration of metabolomics in heart disease and diabetes research: Current achievements and future outlook. Bioanalysis 2011, 3, 2205–2222, doi:10.4155/bio.11.223.

17. Li, M.; Zhou, Z.; Nie, H.; Bai, Y.; Liu, H. Recent advances of chromatography and mass spectrometry in lipidsomics. Anal. Bioanal. Chem. 2011, 399, 243–249, doi:10.1007/s00216-010-3427-7.

18. Courant, F.; Antignac, J.P.; Dervilly-Pinel, G.; Le Bizec, B. Basics of mass spectrometry based metabolomics. Proteomics 2014, 14, 2369–2388, doi:10.1002/pmic.201400255.

19. Cheng, S.; Shah, S.H.; Corwin, E.J.; Fiehn, O.; Fitz Gerald, R. L.; Gerstzen, R. E.; Illig, T.; Rhee, E. P.; Srinivas, P. R.; Wang, T. J.; et al. Potential impact and study considerations of metabolomics in cardiovascular health and disease: A scientiﬁc statement from the American Heart Association. Circ. Cardiovasc. Genet. 2010, 3, e000032, doi:10.1161/CIRCGENETICS.109.940002.

20. Van der Greef, J.; Smilde, A.K. Symbiosis of chemometrics and metabolomics: Past, present, and future. J. Chem. 2015, 39, 1873–1878, doi:10.1002/cem.941.

21. Pauling, L. Orthomolecular psychiatry. Varying the concentrations of substances normally present in the human body may control mental disease. Science 1968, 160, 265–271, doi:10.1126/science.160.3825.265.

22. Pauling, L.; Robinson, A.B.; Teranishi, R.; Cary, P. Quantitative analysis of urine vapor and breath by gas-liquid partition chromatography. Proc. Natl. Acad. Sci. USA 1971, 68, 2374–2376, doi:10.1073/pnas.68.10.2374.

23. Robinson, A.B.; Westall, F.C.; Ellison, G.W. Multiple sclerosis: Urinary amine measurement for orthomolecular diagnosis. Life Sci. 1974, 14, 1747–1753, doi:10.1016/0024-3205(74)90226-8.

24. Horning, E.C.; Horning, M.G. Metabolic proﬁles: Gas-phase methods for analysis of metabolites. Clin. Chem. 1971, 17, 802–809.

25. Horning, E.C.; Horning, M.G. Human metabolic proﬁles obtained by GC and GC/MS. J. Chromatogr. Sci. 1971, 9, 129–140, doi:10.1093/chromsci/9.3.129.

26. Oliver, S.G.; Winson, M.K.; Bell, D.B.; Baganz, F. Systematic functional analysis of the yeast genome. Trends Biotechnol. 1998, 16, 373–378, doi:10.1016/s0167-7799(98)01214-1.

27. Burla, B.; Arita, M.; Arita, M.; Bendt, A. K.; Cazenave-Gassiot, A.; Dennis, E. A.; Ekroos, K.; Han, X.; Ikeda, K.; Liebisch, G.; et al. MS-based lipidomics of human blood plasma: A community-initiated position paper to develop accepted guidelines. J. Lipid Res. 2018, 59, 2001–2017, doi:10.1194/jlr.R087163.

28. Pouraliall Amiri, M.; et al. Metabolomics in early detection and prognosis of acute coronary syndrome. Clin. Chim. Acta 2019, 495, 43–53, doi:10.1016/j.cca.2019.03.1632.

29. Gadian, D. G.; Howitz, D. I.; Radda, G. K.; Seeley, P. J.; Chance, B.; Barlow, C. Phosphorus nuclear magnetic resonance studies on normoxic and ischemic cardiac tissue. Proc. Natl. Acad. Sci. USA 1976, 73, 4446–4448, doi:10.1073/pnas.73.12.4446.

30. Ackerman, J.J.; Bore, P.J.; Gadian, D.G.; Dove, T.H.; Radda, G.K. Nmr studies of metabolism in perfused organs. Philos. Trans. R. Soc. Lond. B Biol. Sci. 1980, 289, 425–436, doi:10.1098/rstb.1980.0059.

31. Bailey, I.A.; Williams, S.R.; Radda, G.K.; Gadian, D.G. Activity of phosphorylase in total global ischaemia in the rat heart. A phosphorus-31 nuclear-magnetic-resonance study. Biochem. J. 1981, 196, 171–178, doi:10.1042/bj1960171.

32. Vignoli, A.; Tenori, L.; Giusti, B.; Valente, S.; Carabba, N.; Balzi, D.; Barchielli, A.; Marchionni, N.; Gensini, G. F.; Marucci, R.; Gori; et al. Differential network analysis reveals metabolic determinants associated with mortality in acute myocardial infarction patients and suggests potential mechanisms underlying different clinical scores used to predict death. Journal of Proteome Res. 2020, 19, 949–961, doi:10.1021/acs.jproteome.9b00779.

33. Yao, W.; Gao, Y.; Wan, Z. Serum metabolomics proﬁling to identify biomarkers for unstable angina. BioMed Res. Int. 2017, 2017, 7657306, doi:10.1155/2017/7657306.
34. Ameta, K.; Gupta, A.; Ameta, D.; Sethi, R.; Kumar, D.; Ahmad, I.; Mahdi, A. A. 1H NMR-derived metabolomics of filtered serum of myocardial ischemia in unstable angina patients. *Clin. Chim. Acta* 2016, 456, 56–62, doi:10.1016/j.cca.2016.02.020.

35. PouralijanAmiri, M.; Khoshkam, M.; Madadi, R.; Kamali, K.; Faghnazadeh Ganji, G.; Salek, R.; Ramazani, A. NMR-based plasma metabolic profiling in patients with unstable angina. *Iran. J. Basic Med. Sci.* 2020, 23, 311–320, doi:10.22038/ijbms.2020.39979.9475.

36. Li, Z.; Liu, X.; Wang, J.; Gao, J.; Guo, S.; Gao, K.; Man, H.; Wang, Y.; Chen, J.; Wang, W. Analysis of urinary metabolic profiling for unstable angina pectoris disease based on nuclear magnetic resonance spectroscopy. *Mol. Biosyst.* 2015, 11, 3387–3396, doi:10.1039/c5mb00489f.

37. Takis, P. G.; Ghini, V.; Tenori, L.; Turano, P.; Luchinat, C. Uniqueness of the NMR approach to metabolomics. *TrAC Trends Anal. Chem.* 2019, 120, 115300, doi:10.1016/j.trac.2018.10.036.

38. Emwas, A.-H.; Roy, R.; McKay, R. T.; Tenori, L.; Sassenti, E.; Gowda, G. A. N.; Raftery, D.; Alahmari, F.; Jaremko, L.; Jaremko, M.; et al. NMR spectroscopy for metabolomics research. *Metabolites* 2019, 9, 123.

39. Senn, T.; Hazen, S. L.; Tang, W. H. Translating metabolomics to cardiovascular biomarkers. *Prog. Cardiovasc. Dis.* 2012, 55, 70–76, doi:10.1016/j.pcd.2012.06.004.

40. Brindle, J. T.; Antti, H.; Holmes, E.; Tranter, G.; Nicholson, J. K.; Bethell, H. W.; Clarke, S.; Schofield, P. M.; McKilligin, E.; Mosedale, D. E.; et al. Rapid and noninvasive diagnosis of the presence and severity of coronary heart disease using 1H-NMR based metabolomics. *Nat. Med.* 2002, 8 (12), 1439–1444.

41. Deng, D.; Liu, L.; Xu, G.; Gan, J.; Shen, Y.; Shi, Y.; Zhu, R.; Lin, Y. Epidemiology and serum metabolic characteristics of acute myocardial infarction patients in chest pain centers. *Iran. J. Public Health* 2018, 47, 1017–1029.

42. Deidda, M.; Piras, C.; Binaghi, G.; Congia, D.; Pani, A.; Boi, A.; Sanna, F.; Rossi, A.; Loi, B.; Cadeddu Dessalvi, C.; et al. Metabolomic fingerprint of coronary artery STEMI patients depends on the ischemic time and inflammatory state. *Sci. Rep.* 2019, 9, 312, doi:10.1038/s41598-018-36415-y.

43. Zhong, W.; Deng, Q.; Deng, X.; Zhong, Z.; Hou, J. Plasma Metabolomics of Acute Coronary Syndrome Patients Based on Untargeted Liquid Chromatography–Mass Spectrometry. *Frontiers in Cardiovascular Medicine* 2021, 8 (277), doi:10.3389/fcvm.2021.616081.

44. Chorell, E.; Olsson, T.; Jansson, J. H.; Wennberg, P., Lyosphospholipids as Predictive Markers of ST-Elevation Myocardial Infarction (STEMI) and Non-ST-Elevation Myocardial Infarction (NSTEMI). *Metabolites* 2020, 11 (1), doi:10.3390/metabo11010025.

45. Aa, N.; Lu, Y.; Yu, M.; Tang, H.; Lu, Z.; Sun, R.; Wang, L.; Li, C.; Yang, Z.; Aa, J.; et al. Plasma Metabolites Alert Patients With Chest Pain to Occurrence of Myocardial Infarction. *Frontiers in Cardiovascular Medicine* 2021, 8 (276), doi:10.3389/fcvm.2021.652746.

46. Chen, H.; Wang, Z.; Qin, M.; Zhang, B.; Lin, L.; Ma, Q.; Liu, C.; Chen, X.; Li, H.; Lai, W.; et al. Comprehensive Metabolomics Identified the Prominent Role of Glycerophospholipid Metabolism in Coronary Artery Disease Progression. *Frontiers in Molecular Biosciences* 2021, 8 (110), doi:10.3389/fmolb.2021.629250.

47. Mehta, A.; Liu, C.; Nayak, A.; Tahhan, A. S.; Ko, Y. A.; Dhindsa, D. S.; Kim, J. H.; Hayek, S. S.; Sperling, L. S.; Mehta, P. K.; et al. Untargeted high-resolution plasma metabolic profiling predicts outcomes in patients with coronary artery disease. *PLoS One* 2020, 15 (8), e0237579, doi:10.1371/journal.pone.0237579.

48. Li, J.; Duan, W.; Wang, L.; Lu, Y.; Shi, Z.; Lu, T. Metabolomics Study Revealing the Potential Risk and Predictive Value of Fragmented QRS for Acute Myocardial Infarction. *J. Proteome Res.* 2020, 19 (8), 3386–3395, doi:10.1021/acs.jproteome.0c00247.

49. Jiang, H.; Jiang, H.; Zhang, J.; Chen, W.; Luo, C.; Li, H.; Hau, W.; Chen, B.; Wang, S., The Serum Metabolic Biomarkers in Early Diagnosis and Risk Stratification of Acute Coronary Syndrome. *Front. Physiol.* 2020, 11, 776–776, doi:10.3389/fphys.2020.00776.

50. Khan, A.; Choi, Y.; Back, J. H.; Lee, S.; Jee, S. H.; Park, Y. H. High-resolution metabolomics study revealing l-homocysteine sulfonic acid, cysteic acid, and carnitine as novel biomarkers for high acute myocardial infarction risk. *Metabolism* 2020, 104, 154051, doi:10.1016/j.metabol.2019.154051.

51. Gundogdu, G.; Senol, O.; Demirkaya Miloglu, F.; Koza, Y.; Gundogdu, F.; Hacimüftüoğlu, A.; Abd El-Aty, A. M. Serum metabolite profiling of ST-segment elevation myocardial infarction using liquid chromatography quadrupole time-of-flight mass spectrometry. *Biomol. Chromatogr.* 2020, 34 (2), e4738, doi:10.1002/bmc.4738.

52. Surendran, A.; Alliani, M.; Ravandi, A. Metabolic characterization of myocardial ischemia-reperfusion injury in ST-segment elevation myocardial infarction patients undergoing percutaneous coronary intervention. *Scientific Reports* 2019, 9 (1), 11742, doi:10.1038/s41598-019-48227-9.

53. Wang, J.; Xu, W.; Zhao, H.; Chen, J.; Zhu, B.; Li, X.; Deng, D.; Wang, J.; Liu, J.; Yu, Y.; et al. Identification of potential plasma biomarkers and metabolic dysfunction for unstable angina pectoris and its complication based on global metabolomics. *Biosci Rep.* 2019, 39 (3), BSR20181658, doi: 10.1042/BSR20181658.

54. Vignoli, A.; Tenori, L.; Giusti, B.; Takis, P. G.; Valente, S.; Ciarabba, N.; Balzi, D.; Barchielli, A.; Marchionni, N.; Gensini, G. F. et al. NMR-based metabolomics identifies patients at high risk of death within two years after acute myocardial infarction in the AMI-Florence II cohort. *BMC Medicine* 2019, 17 (1), 3, doi:10.1186/s12916-018-1240-2.

55. Goulart, V. A. M.; Santos, A. K.; Sandrim, V. C.; Batista, J. M.; Pinto, M. C. X.; Cameron, L. C.; Resende, R. R. Metabolic Disturbances Identified in Plasma Samples from ST-Segment Elevation Myocardial Infarction Patients. *Dis Markers* 2019, 2019, 7676189, doi:10.1155/2019/7676189.

56. Wang, Y.; Sun, W.; Zheng, J.; Xu, C.; Wang, X.; Li, T.; Tang, Y.; Li, Z. Urinary metabolomic study of patients with acute coronary syndrome using UPLC-QTOF/MS. *J. Chromatogr B Analyt Technol Biomed. Life Sci.* 2018, 1100-1101, 122-130, doi:10.1016/j.jchromb.2018.10.005.
57. Du, X.; You, H.; Li, Y.; Wang, Y.; Hui, P.; Qiao, B.; Lu, J.; Zhang, W.; Zhou, S.; Zheng, Y. et al. Relationships between circulating branched chain amino acid concentrations and risk of adverse cardiovascular events in patients with STEMI treated with PCI. *Scientific Reports* 2018, 8 (1), 15809, doi: 10.1038/s41598-018-20198-w.

58. Du, X.; Li, Y.; Wang, Y.; You, H.; Hui, P.; Zheng, Y.; Du, J. Increased branched-chain amino acid levels are associated with long-term adverse cardiovascular events in patients with STEMI and acute heart failure. *Life Sciences* 2018, 209, 167–172, doi: 10.1016/j.lfs.2018.08.011.

59. Huang, L.; Zhang, L.; Li, T.; Liu, Y.-w.; Wang, Y.; Liu, B.-j. Human Plasma Metabolomics Implicates Modified 9-cis-Retinoic Acid in the Phenotype of Left Main Artery Lesions in Acute ST-Segment Elevated Myocardial Infarction. *Scientific Reports* 2018, 8 (1), 12958, doi:10.1038/s41598-018-30219-w.

60. Kohlhauer, M.; Dawkins, S.; Costa, A. S. H.; Lee, R.; Young, T.; Pell, V. R.; Choudhury, R. P.; Banning, A. P.; Kharbanda, R. K.; Saeb-Parsi, K.; et al. Metabolic Profiling in Acute ST-Segment-Elevation Myocardial Infarction Identifies Succinate as an Early Marker of Human Ischemia-Reperfusion Injury. *J. Am. Heart Assoc.* 2018, 7 (8), doi:10.1161/JAHA.117.007546.

61. Zhang, L.; Wei, T. T.; Li, Y.; Li, J.; Fan, Y.; Huang, F. Q.; Cai, Y. Y.; Ma, G.; Liu, J. F.; Chen, Q. Q.; et al. Functional Metabolomics Characterizes a Key Role for N-Acetylnorleucaminic Acid in Coronary Artery Diseases. *Circulation* 2018, 137 (13), 1374-1390, doi:10.1161/CIRCULATIONAHA.117.031139.

62. Yin, X.; de Carvalho, L. P.; Chan, M. Y.; Li, S. F. Y. Integrated metabolomics and metabolomics analyses in acute coronary syndrome patients. *Metabolites* 2017, 9 (6), 734–743, doi:10.3390/m17060071.e.

63. Ali, S. E.; Farag, M. A.; Holvoet, P.; Hanafi, R. S.; Gad, M. Z. A Comparative Metabolomics Approach Reveals Early Biomarkers for Metabolic Response to Acute Myocardial Infarction. *Scientific Reports* 2016, 6 (1), 36359, doi:10.1038/srep36359.

64. Fan, Y.; Li, Y.; Chen, Y.; Zhao, Y. J.; Li, J. L.; Wang, S. L.; Aologa, R. N.; Yin, Y.; Wang, X. M.; et al. Comprehensive Metabolic Characterization of Coronary Artery Diseases. *J. Am. Coll Cardiol* 2016, 68 (12), 1281-93, doi:10.1016/j.jacc.2017.04.032.

65. Xu, X.; Gao, B.; Guan, Q.; Zhang, D.; Ye, X.; Zhou, L.; Tong, G.; Li, H.; Zhang, L.; Tian, J.; et al. Metabolic profile for the early detection of coronary artery disease by using UPLC-QTOF/MS. *J. Pharm Biomed. Anal.* 2016, 129, 34–42, doi:10.1016/j.jpb.2016.06.040.

66. Huang, L.; Li, T. Liu, Y. W.; Zhang, L.; Dong, Z. H.; Liu, S. Y.; Gao, Y. T. Plasma Metabolic Profile Determination in Young ST-segment Elevation Myocardial Infarction Patients with Ischemia and Reperfusion: Ultra-performance Liquid Chromatography Mass Spectrometry for Pathway Analysis. *Sci. Rep.* 2016, 6, doi:10.1038/srep36359.

67. Naz, S.; Calderón Á. A.; García, A.; Gallafrio, J.; Mestre, R. T.; González, E. G.; de Cabo, C. M.; Delgado, M. C. M.; Balanza JÁ, L.; Simionato, A. V. C.; et al Unveiling differences between patients with acute coronary syndrome with and without ST elevation through fingerprinting with CE-MS and HILIC-MS targeted analysis. *Electrophoresis* 2015, 36 (18), 2303-2313, doi:10.1002/elps.201500169.

68. Laborde, C. M.; Mourino-Alvarez, L.; Posada-Ayala, M.; Alvarez-Llamas, G.; Serranillos-Reus, M. G.; Moreu, J.; Vivanco, F.; Patial, D. R.; Barderas, M. G. Plasma metabolomics reveals a potential panel of biomarkers for early diagnosis in acute coronary syndrome. *Metabolites* 2014, 4 (3), 414-424, doi:10.3390/m140300959.s.

69. Sun, M.; Gao, X.; Zhang, D.; Ke, C.; Hou, Y.; Fan, L.; Zhang, R.; Liu, H.; Li, K.; Yu, B. Identification of biomarkers for unstable angina by plasma metabolic profiling. *Mol. Biosyst.* 2013, 9 (12), 3059–3067, doi:10.1039/c3mb70216b.

70. Teul, J.; Garcia, A.; Turién, J.; Martin-Ventura, J. L.; Tarín, N.; Bescós, L. L.; Egido, J.; Barbas, C.; Rupérez, F. J. Targeted and non-targeted metabolic time trajectory in plasma of patients after acute coronary syndrome. *J. Pharm Biomed. Anal.* 2011, 56 (2), 343–351, doi:10.1016/j.jpba.2011.05.020.

71. Vallejo, M.; García, A.; Turión, J.; García-Martínez, D.; Angulo, S.; Martin-Ventura, J. L.; Blanco-Colio, L. M.; Almeida, P.; Egido, J.; Barbas, C.; Plasma fingerprinting with GC-MS in acute coronary syndrome. *Anal. Bioanal. Chem* 2009, 394 (6), 1517–1524, doi:10.1007/s00216-009-9899-5.

72. Griffiths, W. J.; Wang, Y. Mass spectrometry: from proteomics to metabolomics and lipidomics. *Chem Soc. Rev.* 2009, 38 (7), 1882–1896, doi:10.1039/b81555n.

73. Griffiths, J., A Brief History of Mass Spectrometry. *Analytical Chemistry* 2008, 80 (15), 5678-5683, doi:10.1021/ac8013065.

74. Haag, A. M., Mass Analyzers and Mass Spectrometers. In *Modern Proteomics – Sample Preparation, Analysis and Practical Applications*, Mirzaei, H.; Carrasco, M., Eds. Springer International Publishing: Cham, 2016; pp 157 -169, doi:10.1007/978-3-319-41448-5_7.

75. Siuzdak, G., An Introduction to Mass Spectrometry Ionization: An Excerpt from The Expanding Role of Mass Spectrometry in Biotechnology, 2nd ed.; MCC Press: San Diego, 2005. *JALA: Journal of the Association for Laboratory Automation* 2004, 9 (2), 50-63, doi:10.1016/j.jala.2004.01.004.

76. Wilm, M. Principles of electrospray ionization. *Anal. Bioanal Chem* 2011, 407 (7), M111.009407, doi: 10.1002/mcp.M111.009407.

77. Peterson, D. S. Matrix-free methods for laser desorption/ionization mass spectrometry. *Mass Spectrom Rev.* 2007, 26 (1), 19-34, doi:10.1002/mas.20104.

78. Nordström, A.; Want, E.; Northen, T.; Lehtiö, J.; Siuzdak, G. Multiple ionization mass spectrometry strategy used to reveal the complexity of metabolomics. *Anal. Chem.* 2008, 80 (2), 421–429, doi:10.1021/ac701982e.

79. Gowda, G. A.; Djukovic, D. Overview of mass spectrometry-based metabolomics: opportunities and challenges. *Methods Mol. Biol.* 2014, 1198, 3–12, doi:10.1007/978-1-4939-1258-2_1.
101. Au, A. Metabolomics and Lipidomics of Ischemic Stroke. Adv. Clin. Chem 2018, 85, 31–69, doi:10.1016/bs.acc.2018.02.002.

102. Dunn, W. B.; Broadhurst, D. I.; Atherton, H. J.; Goodacre, R.; Griffin, J. L., Systems level studies of mammalian metabolomes: the roles of mass spectrometry and nuclear magnetic resonance spectroscopy. Chem Soc. Rev. 2011, 40 (1), 387-426, doi:10.1039/b906712b.

103. Wishart, D. S., Metabolomics for Investigating Physiological and Pathophysiological Processes. Physiol Rev. 2019, 99 (4), 1819-1875, doi:10.1152/physrev.00035.2018.

104. Stevens, V. L.; Hoover, E.; Wang, Y.; Zanetti, K. A., Pre-Analytical Factors that Affect Metabolite Stability in Human Urine, Plasma, and Serum: A Review. Metabolites 2019, 9 (8), doi:10.3390/metabo9080156.

105. Kamlage, B.; Maldonado, S. G.; Bethan, B.; Peter, E.; Schmitz, O.; Liebenberg, V.; Schatz, P., Quality markers addressing preanalytical variations of blood and plasma processing identified by broad and targeted metabolite profiling. Clin. Chem 2014, 60 (2), 399-412, doi:10.1373/chlm.2013.211979.

106. Liu, X.; Yin, P.; Shao, Y.; Wang, Z.; Wang, B.; Lehmann, R.; Xu, G., Which is the urine sample material of choice for metabolomics-driven biomarker studies? Anal. Chim Acta 2020, 1105, 120–127, doi:10.1016/j.aca.2020.01.028.

107. Giskeødegård, G. F.; Andreassen, T.; Bertilsson, H.; Tessem, M. B.; Batthen, T. F., The effect of sampling procedures and day-to-day variations in metabolomics studies of biofluids. Anal. Chim Acta 2019, 1081, 93–102, doi:10.1016/j.aca.2019.07.026.

108. Van de Merbel, N. C., Quantitative determination of endogenous compounds in biological samples using chromatographic techniques. TrAC Trends in Analytical Chemistry 2008, 27 (10), 924-933, doi:10.1016/j.trac.2008.09.002.

109. Siegel, D.; Permentier, H.; Reijngoud, D. J.; Bischoff, R., Chemical and technical challenges in the analysis of central carbon metabolites by liquid-chromatography mass spectrometry. J. Chromatogr B Analyt Technol Biomed. Life Sci 2014, 966, 21–33, doi:10.1016/j.jchromb.2013.11.022.

110. Zivkovic, A. M.; Wiest, M. M.; Nguyen, U. T.; Davis, R.; Watkins, S. M.; German, J. B., Effects of sample handling and storage on quantitative lipid analysis in human serum. Metabolomics 2009, 5 (4), 507-516, doi:10.1007/s11306-009-0174-2.

111. Jobard, E.; Trédan, O.; Postoly, D.; André, F.; Martin, A. L.; Elena-Herrmann, B.; Boyault, S., A Systematic Evaluation of Blood Serum and Plasma Pre-Analytics for Metabolomics Cohort Studies. Int J. Mol. Sci. 2016, 17 (12), doi:10.3390/jmss17122035.

112. Vuckovic, D., Current trends and challenges in sample preparation for global metabolomics using liquid chromatography-mass spectrometry. Anal. Bioanal Chem 2012, 403 (6), 1523–1548, 10.1007/s00216-012-0639-y.

113. Bowen, R. A.; Chan, Y.; Cohen, J.; Rebak, N. N.; Hortin, G. L.; Csako, G.; Remaley, A. T., Effect of blood collection tubes on total triiodothyronine and other laboratory assays. Clin. Chem 2005, 51 (2), 424–433, doi:10.1373/chlm.2004.043349.

114. Clark, S.; Youngman, L. D.; Palmer, A.; Parish, S.; Peto, R.; Collins, R., Stability of plasma analytes after delayed separation of whole blood: implications for epidemiological studies. Int J. Epidemiol. 2003, 32 (1), 125–130, doi:10.1093/ije/dyg023.

115. Teahan, O.; Gamble, S.; Holmes, E.; Waxman, J.; Nicholsoon, J. K.; Bevan, C.; Keun, H. C., Impact of analytical bias in metabolomic studies of human blood serum and plasma. Anal. Chem. 2006, 78 (13), 4307–4318, doi:10.1021/ac051972y.

116. Liu, X.; Hoene, M.; Wang, X.; Yin, P.; Häring, H. U.; Xu, G.; Lehmann, R., Serum or plasma, what is the difference? Investigations to facilitate the sample material selection decision making process for metabolomics studies and beyond. Anal. Chim Acta 2018, 1037, 295–300, doi:10.1016/j.aca.2018.03.009.

117. Wedge, D. C.; Allwood, J. W.; Dunn, W.; Vaughan, A. A.; Simpson, K.; Brown, M.; Priest, L.; Blackhall, F. H.; Whetton, A. D.; Dive, C.; et al. Is serum or plasma more appropriate for intersubject comparisons in metabolomic studies? An assessment in patients with small-cell lung cancer. Anal. Chem. 2011, 83 (17), 6689–6697, doi:10.1021/ac112224.

118. Yu, Z.; Kastenmüller, G.; He, Y.; Belcredì, P.; Möller, G.; Frehn, C.; Mendes, J.; Wahl, S.; Roemisch-Margl, W.; Ceglarek, U.; et al. Differences between human plasma and serum metabolite profiles. PLoS One 2011, 6 (7), e21230, doi:10.1371/journal.pone.0021016.

119. Malik, D. M. Rhoades, S.; Weljie, A., Extraction and Analysis of Pan-metabolome Polar Metabolites by Ultra Performance Liquid Chromatography-Tandem Mass Spectrometry (UPLC-MS/MS). Bio Protoc. 2018, 8 (3), e2715, doi:10.21769/BioProtoc.2715.

120. Tambellini, N. P.; Zaremberg, V.; Turner, R. J.; Weljie, A. M., Evaluation of extraction protocols for simultaneous polar and non-polar yeast metabolite analysis using multivariate projection methods. Metabolites 2013, 3 (3), 592-605, doi:10.3390/metabo3030592.

121. Hyötyniemi, T., Critical evaluation of sample pretreatment techniques. Anal. Bioanal Chem 2009, 394 (3), 743–758, doi:10.1007/s00216-009-2772-2.

122. Panuwet, P.; Hunter, R. E., Jr.; D’Souza, P. E.; Chen, X.; Radford, S. A.; Cohen, J. R.; Marder, M. E.; Kartvenka, K.; Ryan, P. B.; Barr, D. B., Biological Matrix Effects in Quantitative Tandem Mass Spectrometry-Based Analytical Methods: Advancing Bio-monitoring. Crit Rev. Anal. Chem. 2016, 46 (2), 93-105, doi:10.1080/10408347.2014.980775.

123. Tsakelidou, E.; Virgiliou, C.; Valianou, L.; Gika, H. G.; Raikos, N.; Theodoridis, G., Sample Preparation Strategies for the Effective Quantitation of Hydrophilic Metabolites in Serum by Multi-Targeted HILIC-MS/MS. Metabolites 2017, 7 (2), doi:10.3390/metabo7020013.

124. Kole, P. L.; Venkatesh, G.; Kotecha, J.; Sheshala, R., Recent advances in sample preparation techniques for effective bioanalytical methods. Biomed. Chromatogr. 2011, 25 (1-2), 199-217, doi:10.1002/bmc.1560.

125. Alshammari, T. M.; Al-Hassan, A. A.; Hadda, T. B.; Aljofan, M., Comparison of different serum sample extraction methods and their suitability for mass spectrometry analysis. Saudi Pharm J. 2015, 23 (6), 689–697, doi:10.1016/j.jspj.2015.01.023.

126. Rico, E.; González, O.; Blanco, M. E.; Alonso, R. M., Evaluation of human plasma sample preparation protocols for untargeted metabolic profiles analyzed by UHPLC-ESI-TOF-MS. Anal. Bioanal Chem 2014, 406 (29), 7641–7652, doi:10.1007/s00216-014-8212-y.
127. Heather, L. C.; Wang, X.; West, J. A.; Griffin, J. L., A practical guide to metabolomic profiling as a discovery tool for human heart disease. *J. Mol. Cell Cardiol* 2013, 55, 2–11, doi:10.1016/j.yjmcc.2012.12.001.

128. Chang, M. S.; Ji, Q.; Zhang, J.; El-Shourbagy, T. A., Historical review of sample preparation for chromatographic bioanalysis: pros and cons. *Drug Development Research* 2007, 68 (3), 107-133, doi:10.1016/d/dr.2007.12.001.

129. Biddlecombe, R. A.; Pleasance, S., Automated protein precipitation by filtration in the 96-well format. *J. Chromatogr B Biomed. Sci Appl* 1999, 734 (2), 257–265, doi:10.1016/S0378-4347(99)00355-2.

130. Ma, J.; Shi, J.; Le, H.; Cho, R.; Huang, J. C.; Miao, S.; Wong, B. K., A fully automated plasma protein precipitation sample preparation method for LC-MS/MS bioanalysis. *J. Chromatogr B Analyt Technol Biomed. Life Sci* 2008, 862 (1-2), 219–226, doi:10.1016/j.jchromb.2007.12.012.

131. Li, M.; Yang, L.; Bai, Y.; Liu, H., Analysis methods in lipidomics and their applications. *Anal. Chem* 2014, 86 (1), 161–175, doi:10.1021/ac403554h.

132. Folch, J.; Lees, M.; Sloane Stanley, G. H., A simple method for the isolation and purification of total lipides from animal tissues. *J. Biol Chem* 1957, 226 (1), 497-509, doi:10.1010/jb.1957.v226.11.09-509.

133. Matyash, V.; Liebisch, G.; Kurzchalia, T. V.; Shevchenko, A.; Schwudke, D., Lipid extraction by methyl-tert-butyl ether for high-throughput lipidomics. *J. Lipid Res.* 2008, 49 (5), 1137–1146, doi:10.1194/jlr.D700041-JLR200.

134. Löfgren, L.; Ståhlin, M.; Forsberg, G. B.; Saarinen, S.; Nilsson, R.; Hansson, G. L., The BUME method: a novel automated chloroform-free 96-well total lipid extraction method for blood plasma. *J. Lipid Res.* 2012, 53 (8), 1690–1700, doi:10.1194/jlr.D023036.

135. Barnes, B. B.; Snow, N. H., 3.44 - Recent Advances in Sample Preparation for Explosives. In *Comprehensive Sampling and Sample Preparation*, Pawliszyn, J., Ed. Academic Press: Oxford, 2012, pp 893-926, doi:10.1016/S0166-526X(02)80049-6.

136. Sitnikov, D. G.; Mornin, C. S.; Vuckovic, D., Systematic Assessment of Seven Solvent and Solid-Phase Extraction Methods for Metabolomics Analysis of Human Plasma by LC-MS. *Scientific Reports* 2016, 6 (1), 38885, doi:10.1038/srep38885.

137. Michopoulos, F.; Lai, L.; Gika, H.; Theodordinis, G.; Wilson, I., UPLC-MS-Based Analysis of Human Plasma for Metabolomics Using Solvent Precipitation or Solid Phase Extraction. *Journal of Proteome Research* 2009, 8 (4), 2114-2121, doi:10.1021/pr801045q.

138. Kordalewska, M.; Markuszewski, M. J., Metabolomics in cardiovascular diseases. *Journal of Pharmaceutical and Biomedical Analysis* 2015, 113, 121–136, doi:10.1016/j.jpba.2015.04.021.

139. Xia, J.; Psychogios, N.; Young, N.; Wishart, D. S., MetaboAnalyst: a web server for metabolomic data analysis and interpretation. *Nucleic Acids Res.* 2009, 37 (Web Server issue), W652-W660, doi:10.1093/nar/gkp356.

140. Forsberg, E. M.; Huan, T.; Rineland, D.; Benton, H. P.; Warth, B.; Hilmers, B.; Stuzdak, G., Data processing, multi-omic pathway mapping, and metabolite activity analysis using XCMS Online. *Nat. Protoc* 2018, 13 (4), 633-651, doi:10.1038/nprot.2017.151.

141. Zhang, X.; Dong, J.; Raftery, D., Five Easy Metrics of Data Quality for LC-MS-Based Global Metabolomics. *Analytical chemistry* 2020, 92 (19), 12925-12933, doi:10.1021/acs.analchem.0c01493.

142. Livera, A. M. D.; Sysi-Aho, M.; Jacob, L.; Folch, J.; Le, H.; Cho, R.; Huang, J. C.; Miao, S.; Wong, B. K., A fully automated plasma protein precipitation sample preparation method for LC-MS/MS bioanalysis. *J. Chromatogr B* 862, 219–226, doi:10.1016/j.jchromb.2007.12.012.

143. Sabatine, M. S.; Liu, E.; Morrow, D. A.; Heller, E.; McCarroll, R.; Wiegand, R.; Berriz, G. F.; Roth, F. P.; Gerszten, R. E., Metabolomic Identification of Novel Biomarkers of Myocardial Ischemia. *Circulation* 2005, 112 (25), 3868-3875, doi:10.1161/CIRCULATIONAHA.105.569137.

144. Lewis, G. D.; Wei, R.; Liu, E.; Yang, E.; Shi, X.; Martinovic, M.; Farrell, L.; Asnani, A.; Cyrille, M.; Ramanathan, A.; et al. Metabolite profiling of blood from individuals undergoing planned myocardial infarction reveals early markers of myocardial injury. *J. Clin. Invest.* 2008, 118 (10), 3503–3512, doi:10.1172/JCI35111.
153. Cao, B.; Wang, D.; Pan, Z.; Brietke, E.; McIntyre, R. S.; Musial, N.; Mansur, R. B.; Subramaniapillai, M.; Zeng, J.; Huang, N.; et al. Characterizing acyl-carnitine biosignatures for schizophrenia: a longitudinal pre- and post-treatment study. *Transl Psychiatry* 2019, 9 (1), 19-19, doi:10.1038/s41398-018-01353-x.

154. Diamanti, K.; Cavalli, M.; Pan, G.; Pereira, M. J.; Kumar, C.; Skritic, S.; Grabherr, M.; Risérus, U.; Eriksson, J. W.; Komorowski, J.; et al. Intra- and inter-individual metabolic profiling highlights carnitine and lysophosphatidylcholine pathways as key molecular defects in type 2 diabetes. *Scientific reports* 2019, 9 (1), 9653-9653, doi:10.1038/s41598-019-45906-5.

155. Guasch-Ferré, M.; Zheng, Y.; Ruiz-Canela, M.; Hruby, A.; Martínez-González, M. A.; Clish, C. B.; Corella, D.; Estruch, R.; Ros, E.; Fitó, M.; et al. Plasma acylcarnitines and risk of cardiovascular disease: effect of Mediterranean diet interventions. *Am. J. Clin. Nutr.* 2016, 103 (6), 1408-1416, doi:10.3945/ajcn.116.130492.

156. Makrecka-Kuka, M.; Sevostjanovs, E.; Vilks, K.; Volska, K.; Antone, U.; Kuka, J.; Makarova, E.; Pugovics, O.; Dambrova, M.; Liepins, E., Plasma acylcarnitine concentrations reflect the acylcarnitine profile in cardiac tissues. *Scientific Reports* 2017, 7 (1), 17528, doi:10.1038/s41598-017-17797-x.

157. Law, S.-H.; Chan, M.-L.; Marathe, G. K.; Parveen, F.; Chen, C.-H.; Ke, L.-Y., An Updated Review of Lysophosphatidylcholine Metabolism in Human Diseases. *International journal of molecular sciences* 2019, 20 (5), 1149, doi:10.3390/ijms20051149.

158. Sasagawa, T.; Suzuki, K.; Shiota, T.; Kondo, T.; Okita, M., The significance of plasma lysophospholipids in patients with renal failure on hemodialysis. *J. Nutr Sci Vitamino (Tokyo)* 1998, 44 (6), 809-818, doi:10.3177/jnsv.44.809.

159. Okita, M.; Gaudette, D. C.; Mills, G. B.; Holub, B. J.; Elevated levels and altered fatty acid composition of plasma lysophosphatidylcholine(lysoPC) in ovarian cancer patients. *Int J. Cancer* 1997, 71 (1), 31–34, doi:10.1002/(sici)1097-0215(19970328)71:1<31::aid-ijc7>3.0.co;2-4.

160. Rabin, R. A.; Galassi, R.; Fumelli, P.; Dousset, N.; Solera, M. L.; Valdiguié, P.; Curatola, G.; Ferretti, G.; Taus, M.; Mazzanti, L., Reduced Na(+)+-K(+)+-ATPase activity and plasma lysophosphatidylcholine concentrations in diabetic patients. *Diabetes* 1994, 43 (7), 915-919, doi:10.2337/diab.43.7.915.

161. Tan, S. T.; Ramesh, T.; Toh, X. R.; Nguyen, L. N., Emerging roles of lysophospholipids in health and disease. *Progress in Lipid Research* 2020, 80, 101068, doi:10.1016/j.plipres.2020.101068.

162. Carneiro, A. V.; Costa, J.; Borges, M., Statins for primary and secondary prevention of coronary heart disease. A scientific review. *Rev. Port. Cardiol* 2004, 23 (1), 95-122.

163. Hausenloy, D. J.; Yellon, D. M., Myocardial ischemia-reperfusion injury: a neglected therapeutic target. *J. Clin. Invest.* 2013, 123 (1), 92-100, doi:10.1172/JCI62874.

164. Keeley, E. C.; Boura, J. A.; Grines, C. L., Primary angioplasty versus intravenous thrombolytic therapy for acute myocardial infarction: a quantitative review of 23 randomised trials. *The Lancet* 2003, 361 (9351), 13-20, doi:10.1016/S0140-6736(03)12113-7.

165. Kalogeris, T.; Baines, C. P., Cell biology of ischemia/reperfusion injury. *Int Rev. Cell Mol. Biol* 2015, 238, 229–317, doi:10.1016/B978-0-12-394309-5.00006-7.

166. Chouchani, E. T.; Pell, V. R.; Gaude, E.; Aksentijevic, D.; Sundier, S. Y.; Robb, E. L.; Logan, A.; Nadtochiy, S. M.; Ord, E. N. J.; Smith, A. C.; et al. Ischaemic accumulation of succinate controls reperfusion injury through mitochondrial ROS. *Nature* 2014, 515 (7527), 431-435, doi:10.1038/nature13909.

167. Li, T.; Zhang, Z.; Kolwicz, S. C., Jr.; Abell, L.; Roe, N. D.; Kim, M.; Zhou, B.; Cao, Y.; Ritterhoff, J.; Gu, H.; et al. Defective Branched-Chain Amino Acid Catabolism Disrupts Glucose Metabolism and Sensitizes the Heart to Ischemia-Reperfusion Injury. *Cell Metab* 2017, 25 (2), 375-385, doi:10.1016/j.cmet.2016.11.003.

168. Dennis, E. A.; Norris, P. C., Eicosanoid storm in infection and inflammation. *Nat. Rev. Immunol* 2015, 15 (8), 511-523, doi:10.1038/nri3859.

169. Li-Gao, R.; Hughes, D. A.; le Cessie, S.; de Mutsert, R.; den Heijer, M.; Rosendaal, F. R.; Willems van Dijk, K.; Timpson, N. J.; Mooij-Kanamori, D. O., Assessment of reproducibility and biological variability of fasting and postprandial plasma metabolite concentrations using 1H NMR spectroscopy. *PLOS ONE* 2019, 14 (6), e0218549, doi:10.1371/journal.pone.0218549.

170. Sampson, J. N.; Boca, S. M.; Shu, X. O.; Stolzenberg-Solomon, R. Z.; Matthews, C. E.; Hsing, A. W.; Tan, Y. T.; Ji, B.-T.; Chow, W.-H.; Cai, Q.; et al. Metabolomics in epidemiology: sources of variability in metabolite measurements and implications. *Cancer Epidemiol Biomarkers Preva* 2012, 22 (4), 631-640, doi:10.1158/1055-9965.EPI-12-1109.

171. Johnson, C. H.; Ivanisevic, J.; Siuzdak, G., Metabolomics: beyond biomarkers and towards mechanisms. *Nat. Rev. Mol. Cell Biol* 2016, 17 (7), 451-459, doi:10.1038/nrm.2016.25.

172. Pinu, F. R.; Goldansaz, S. A.; Jaine, J., Translational Metabolomics: Current Challenges and Future Opportunities. *Metabolites* 2019, 9 (6), 108, doi:10.3390/metabo9060108.