Emerging roles of non-histone protein crotonylation in biomedicine

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Abstract
Crotonylation of proteins is a newly found type of post-translational modifications (PTMs) which occurs leadingly on the lysine residue, namely, lysine crotonylation (Kcr). Kcr is conserved and is regulated by a series of enzymes and co-enzymes including lysine crotonyltransferase (writer), lysine decrotonylase (eraser), certain YEATS proteins (reader), and crotonyl-coenzyme A (donor). Histone Kcr has been substantially studied since 2011, but the Kcr of non-histone proteins is just an emerging field since its finding in 2017. Recent advances in the identification and quantification of non-histone protein Kcr by mass spectrometry have increased our understanding of Kcr. In this review, we summarized the main proteomic characteristics of non-histone protein Kcr and discussed its biological functions, including gene transcription, DNA damage response, enzymes regulation, metabolic pathways, cell cycle, and localization of heterochromatin in cells. We further proposed the performance of non-histone protein Kcr in diseases and the prospect of Kcr manipulators as potential therapeutic candidates in the diseases.

Keywords: Lysine crotonylation, Post-translational modification, Protein, Cell biology, Disease

Introduction
Post-translational modifications (PTMs) of proteins make up most of the proteome and to a large extent, and establish the impressive level of functional diversity in higher multi-cellular organisms. Mounting evidence suggests that PTMs provide an elegant mechanism to govern protein function in diverse biological processes including cell differentiation and organismal development, and aberrant protein modification may contribute to diseases such as cancer. As the only amino acid with a side chain amine [1], lysine can be covalently modified by glycosyl [2], propionyl [3], butyryl [4], acetyl [5], hydroxyl [6], crotonyl [7], ubiquitinyl and ubiquitinyl-like [8], formyl [9], malonyl [10], succinyl [11], and methyl [12] groups. Some modifications, such as acetylation, methylation, and ubiquitination, have been extensively studied and are considered to be related to chromatin remodeling, gene transcriptional regulation, and other key biological processes.

Among the above PTMs, lysine crotonylation (Kcr) represents a newly discovered and often elusive PTM that has nonetheless the potential to alter the function of the modified protein. Histone crotonylation was first reported as a special marker of sex-related genes and was directly related to gene transcription [7]. Subsequently, some studies found that lysine crotonylation can play an important role in many diseases such as acute renal injury, depression, HIV latency, and cancer process by regulating the structure and function of histone [13–16]. In the past decade, mass spectrometry (MS)-based proteomics has greatly accelerated the discovery and identification of endogenous crotonylated proteins, and revealed the regulatory process of non-histone crotonylation. The discovery and identification of mammalian lysine crotonyltransferases (KCTs), lysine decrotonylases (KDCRs), and the crotonylation reader domain pave the way for the study of non-histone protein crotonylation.
The identification of thousands of crotonylation sites has arisen great interest of biomedical researchers. More importantly, a growing number of studies have shown that non-histone protein crotonylation is involved in almost all major biological processes [17–21].

Here, we provide an overview of the expanding landscape of non-histone protein crotonylation, including the regulation, large-scale identification, biological function of protein crotonylation, and the crosstalk of Kcr with other PTMs. We further discuss the connection between protein crotonylation and diseases in which Kcr may be potential therapeutic targets of the diseases. Although there are not many reported in-depth investigations on the crotonylation of non-histone proteins, a comprehensive review on this rapidly developing field may help to uncover its biomedical significance and prospects.

Uncovering lysine crotonylation

It is increasingly appreciated that combinations of PTMs can generate distinct protein isoforms with varying functions, which vastly expand the functional diversity of mammalian proteomes. The widespread occurrence of PTMs only started to become clear in the first years of the twenty-first century, when advances in high-resolution mass spectrometry enabled detection of thousands of low-abundance PTM sites. In this context, protein Kcr on histones was first described in 2011 by Zhao and colleagues [7], who designed a comprehensive method to systematically analyze histone PTM, using PTMap, an algorithm that can recognize all possible PTMs of proteins [22]. In this method, mass spectrometry (MS) analysis of histone hydrolyzed peptides maximizes sequence coverage and sensitivity, resulting in recognition of many new PTM sites, including Kcr identified as a new type of histone modification.

Histone crotonylation is an evolutionarily conserved histone post-translational modification appearing in eukaryotic cells from a wide range of species. Using a pan antibody against Kcr, Kcr signals in the core histones of sapiens (HeLa) cells, mouse, cerevisiae, elegans, melanogaster, as well as plant, have been detected [7, 23–26]. The modified proteins and sites vary and overlap among different species, such as H2AK125cr in humans and mice [26].

These groundbreaking discoveries set the stage for the field of non-histone protein crotonylation. In 2017, Xu et al. [17] first reported that in addition to histones, Kcr also occurs on a variety of non-histone proteins. Increasing proteomic analyses have demonstrated that the frequency of non-histone protein crotonylation is very high, and these proteins are involved in many key cellular processes which are related to physiology and disease in eukaryotic cells. These mass spectrometry-based studies have quantified the relative effects of tens of thousands of crotonylation sites on genetic and metabolic activities, and provided insight into the dynamic regulation of Kcr.

Regulation of crotonylation

There are structural similarities between lysine crotonylation and lysine acetylation, and in fact, many of the “modulators” of crotonylation are the same as those of acetylation [27, 28]. However, crotonylation is unique in its π-electron conjugation, resulting in being rigid and planar in shape. Crotonyllysine but not acetyllysine is found to more robustly neutralize the positive charge because of a longer carbon chain [29]. Histone crotonylation have been found sharing the similar set of regulatory machineries as histone acetylation [19, 30]. In the process of crotonylation, crotonyl is added to the substrate proteins by lysine crotonyltransferase (KCT) (“writer”) [17, 31–33] and then histone crotonylation can be recognized by another family of proteins such as YEATS2, AF9, and Taf14 (“reader”) [29, 34, 35], which usually initiates a series of downstream signals [29]. At the end of PTM-induced signal transduction, most crotonylation can also be removed by lysine decrotonylase (KDCR) (“eraser”) [17, 36–38]. Part of the effect of either KCT or KDCR can be selectively counteracted by drug inhibitors (Fig. 1). The crotonyltransferase, decrotonylase, reader, and donor of non-histone and histone crotonylation do not completely overlap, which are respectively described below.

Lysine crotonyltransferase (writer), decrotonylase (eraser), and reader

The modulators of histone crotonylation have been extensively studied and summarized earlier [19, 39], and some of them can also regulate non-histone crotonylation. Six KCTs have been identified in metazoans and are classified into three major families: P300/CBP (p300/CREB-binding protein) [40], MYST (Moz, Ybf2, Sas2, and Tip60) [32], and GNAT (GCN5-related N-acetyltransferase) [33]. So far, P300 and hMOF in MYST family and PCAF in GNAT family were revealed to be non-histone crotonyltransferase (Table 1). P300 and its homologue CBP are two highly related KCTs that function as general transcriptional coactivators. The human males absent on the first (hMOF) protein which belongs to the MYST family, and the P300/CBP-associated factor (PCAF) which is a member of the GNAT family, both possess evolutionarily conserved KCT activity [17, 32]. CBP, p300, and hMOF have the same enzyme reaction centers which catalyze acetylation and crotonylation at multiple sites. It is found that the activity of each KCT varies from substrate to substrate (Table 1). The non-histone NPM1 can be strongly crotonylated by CBP and hMOF, but only moderately crotonylated by PCAF.
However, crotonylation of DDX5 can be catalyzed by CBP, but not by other KCTs [17].

Most lysine decrotonylases have a wide range of substrates, including histone and non-histone proteins. KDCRs can be grouped into two major categories: Zn$^{2+}$-dependent KDCRs and NAD$^+$-dependent sirtuin decrotonylases (Table 2). The Zn$^{2+}$-dependent KDCRs are often referred to as classical HDACs and belong to class I [41]. In class I HDACs, HDAC1 and HDAC3, but not HDAC2, can decrotonylase non-histone NPM1, and

### Table 1 Lysine crotonyltransferases and their histone and non-histone substrates

| Family          | Enzyme name | Substrates and modified lysine residues | Histone | References | Non-histone | References |
|-----------------|-------------|----------------------------------------|---------|------------|-------------|------------|
| p300/CBP family | p300, CBP   | H3K18                                  | [31]    | NPM1, DDXS | [17]        |
| MYST family     | hMOF        | H3K4, H3K9, H3K18, H3K23, H4K8, and H4K12 | [32]    | NPM1       |             |
|                 | Esa1        | H4K5, H4K8, H4K12, and H4K16           | [33]    | N/A        | N/A         |
| GNAT family     | GCN5        | H3K9, K3K14, and H3K18A                | [33]    | N/A        | N/A         |
|                 | PCAF        | N/A                                    |         | N/A        | NPM1 [17]   |

DDX5 DEAD-box helicase 5, Esa1 essential Sas2-related acetyltransferase 1, GCN5 general control nonrepressed-protein 5, GNAT GCN5-related N-acetyltransferase, hMOF human males absent on the first, MYST Moz, Ybf2, Sas2, and Tip60, NPM1 nucleophosmin-1, N/A not available, PCAF P300/CBP-associated factor, P300/CBP P300/CREB-binding protein.
this effect can be reversed by treatment with HDACs inhibitors such as trichostatin A (TSA). Similarly, two other HDAC inhibitors, suberoylanilide hydroxamic acid (SAHA) and LBH589, can promote the crotonylation of NPM1 [17] (Fig. 1). Sirtuin family deacetylases (SIRTs), which are also referred to as class III HDACs, localize to different cellular compartments, including the nucleus (SIRT1), cytoplasm (SIRT2), and mitochondria (SIRT3) [42, 43]. So far, it has not been reported that Sirtuin family can decrotonylate non-histone protein. In the function and regulation of protein crotonylation, Kcr must be recognized by certain proteins possessing special structures, these proteins are called reader, including bromodomain, double plant homeodomain (PHD) finger (DPF), and YEATS domains [29, 34, 35, 44–46]. Reader originally refers to a chromatin binding protein module that recognizes the covalent modification of histone. At present, there is no evidence that non-histone KCR also has a reader.

**Crotonyl-CoA (donor)**
Crotonyl-CoA is presumed a crotonyl donor in the process of Kcr. Crotonate, mainly produced by the colon microbiota, is the short-chain fatty acid (SCFA) precursor of crotonyl-CoA [47]. Circulating short-chain fatty acids (SCFA), such as acetate, crotonate, butyrate, and propionate, are taken up by tissues and converted to acyl-CoA by the acyl-CoA synthetase short chain family member 2 (ACSS2) or eventually yield crotonyl-CoA through different metabolic pathways such as fatty acid β-oxidation pathway and lysine degradation [48] (Fig. 1). Intracellular and intercellular metabolism can affect Kcr by changing the concentration of crotonyl-CoA in the following two ways:

First, direct elevation of crotonyl-CoA by sodium crotonate. Sodium crotonate (NaCr) enhances non-histone crotonylation most likely through conversion of NaCr to crotonyl-CoA. Adding NaCr to the cell culture media can lead to a significant increase in the crotonylation of many non-histone proteins, including ubiquitin ligase RNF2 and the binding enzymes UBE2E1, NCoR1, and RBP4, which are components of histone deacetylase complex 12 [49]. Crotonylation of proteins induced by NaCr may also be catalyzed by enzymes. NaCr can specifically enhance protein crotonylation but not acetylation in wild HeLa cells, and this induction can be nearly completely eliminated by C646 and SGC-CBP30, two selective P300/CBP inhibitors [49].

Second, enzymatic regulation of crotonyl-CoA level. The enzymes involved in the conversion of crotonate to crotonyl-CoA can affect the crotonylation. Chromodomain Y-like (CDYL) protein, which contains an N-terminal chromodomain and a C-terminal enoyl-coenzyme A hydratase/isomerase catalytic domain (also known as CoA pocket or CoAP), has been implicated in epigenetic regulation and transcription repression by targeting chromatin through its N-terminal chromodomain [50]. Liu et al. [51] reported that CDYL, acting as a crotonyl-CoA hydratase, can add a water molecule to the double bond at the position between the second and third carbon, and negatively regulate histone Kcr (Fig. 1). Yu et al. [52] reported that CDYL can negatively regulate the Kcr of RPA1 on multiple lysine sites, which is a reaction to DNA-damaging insults.

**Large-scale identification of non-histone protein crotonylation**
The breakthroughs in high-resolution mass spectrometry (MS)-based proteomics have enabled both the identification of thousands of crotonylation sites and the relative quantification of these sites in a single experiment, ushering in the age of crotonylomics (proteome-wide characteristics of crotonylation). Crotonylomics mapping is a prerequisite for systems-level analyses and provides an important resource for discovering novel properties and regulatory functions of crotonylation.

| Co-factor | Class | Members | Location     | Histone                           | Substrate                                                                 |
|-----------|-------|---------|--------------|-----------------------------------|---------------------------------------------------------------------------|
| Zn^{2+}-dependent | HDAC1 | HDAC1   | Nucleus      | H3K4, H3K9, H3K18, H3K23, H4K8, and H4K12 | NPM1 [17]                                                                 |
|           |       | HDAC2   | Nucleus      | H3K9, H3K18, and H4K8             | N/A N/A                                                                   |
|           |       | HDAC3   | Nucleus      | H3K9, H3K18, and H4K8             | NPM1 [17]                                                                 |
|           |       | HDAC8   | Nucleus      | H3K4, H3K9, H3K18, H4K8, and H4K12 | N/A N/A                                                                   |
| NAD^{+}-dependent | HDAC III | SIRT1  | Nucleus/cytoplasm | H3K4, H3K9, and H4K8 | N/A N/A                                                                   |
|           |       | SIRT2   | Cytoplasm    | H3K4 and H3K9                    | N/A N/A                                                                   |
|           |       | SIRT3   | Mitochondria | H3K4                             | N/A N/A                                                                   |

N/A not available

Table 2 Different classes of lysine decrotonylases, their co-factors, cellular locations, and substrates

| Co-factor | Class | Members | Location | Histone                           | Substrate                                                                 |
|-----------|-------|---------|----------|-----------------------------------|---------------------------------------------------------------------------|
| Zn^{2+}-dependent | HDAC1 | HDAC1   | Nucleus  | H3K4, H3K9, H3K18, H3K23, H4K8, and H4K12 | NPM1 [17]                                                                 |
|           |       | HDAC2   | Nucleus  | H3K9, H3K18, and H4K8             | N/A N/A                                                                   |
|           |       | HDAC3   | Nucleus  | H3K9, H3K18, and H4K8             | NPM1 [17]                                                                 |
|           |       | HDAC8   | Nucleus  | H3K4, H3K9, H3K18, H4K8, and H4K12 | N/A N/A                                                                   |
| NAD^{+}-dependent | HDAC III | SIRT1 | Nucleus/cytoplasm | H3K4, H3K9, and H4K8 | N/A N/A                                                                   |
|           |       | SIRT2   | Cytoplasm | H3K4 and H3K9                    | N/A N/A                                                                   |
|           |       | SIRT3   | Mitochondria | H3K4                             | N/A N/A                                                                   |
Quantitative mass spectrometry for crotonylomics analysis

Various MS instruments and methodological approaches can be used to perform crotonylomics analysis for proteins and peptides. Nearly all large-scale crotonylation studies use the “shotgun” or bottom-up proteomics approach, which involves enzymatic digestion of all proteins (the proteome) followed by liquid chromatography coupled to tandem MS (LC–MS/MS) (Fig. 2).

Crotonylation of some protein sites cannot be detected in wild type cells. In order to find out which lysine sites may be crotonylated, some studies use drugs that promote crotonylation or knock down the expression of “modulators” of crotonylation during cell culture, such as sodium crotonate (NACR) which promotes non-histone crotonylation by conversion to crotonyl-coenzyme A [49], and HDAC inhibitor SAHA [21] or crotonyl-CoA hydrolase CDYL [52] which negatively regulate Kcr. But more researchers collect biological samples under interested conditions for modification (Table 3).

In the next workflow, proteins are extracted from biological samples and digested into large numbers of peptides by enzymatic hydrolysis (usually trypsin), only a small part of the peptides is crotonylated. In order to identify the modified sites under specific conditions, quantitative analysis should be used. The most commonly used methods are to identify modified sites by comparing the intensity of crotonylated peptides among different samples, including metabolic labeling, chemical labeling, and label-free quantification. Each method has its advantages and disadvantages. SILAC (stable isotope labelling by amino acids in cell culture), which belongs to metabolic labeling, is one of the most commonly used methods in quantitative proteomics [53]. Natural isotopes (light) or stable isotopes (medium/heavy) are used to replace the corresponding amino acids during cell culture. Its advantage is the high efficiency for protein labeling, while its disadvantage is time consuming. Metabolic labeling has been successfully used to reveal the relative

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Fig. 2 Schematic diagram of the experimental procedures for mass spectrometry-based global analysis. Proteins are extracted from cells or tissues and digested into peptides with a protease such as trypsin. The tryptic peptides are then separated and fractionated by high pH reverse-phase high performance liquid chromatography (HPLC). Proteolysis of whole-cell protein extracts generates numerous peptides, but only a small fraction is crotonylated. To enrich lysine crotonylated peptides, pan-Kcr antibodies are applied to identify the crotonylated peptides in complex peptide mixtures using immunomass purification. The resulting peptides are ionized in the electrospray source before entering the mass spectrometer. MS and MS/MS spectra are then computationally processed to deduce peptide sequences, including the presence and location of crotonylation, and to quantify the abundance of crotonylated peptides and proteins.
Table 3: Global profiling of non-histone crotonylation based on LC–MS/MS analysis

| Biological samples analyzed | Experiment condition | Identified proteins | Kcr sites | Amino acids flanking the identified Kcr sites | Subcellular localization | Biological function | Validated proteins by WB | Year | References |
|----------------------------|----------------------|--------------------|-----------|-----------------------------------------------|--------------------------|--------------------|-------------------------|------|------------|
| H. sapiens                 | PBMCs were obtained and compared with normal controls | 353                | 770       | Alanine (A), aspartic acid (D), and lysine (K) residues at the −1 to +1 positions | Organelles, extracellular regions and membranes (69%) | Upregulated proteins involved: focal adhesion, cell-substrate junction, cell-substrate adherens junction, biological process of regulation of body fluid levels, and calcium ion binding. Downregulated proteins involved: cellular component of blood microparticle, biological process of humoral immune response, and ubiquitin-like protein ligase binding. | N/A | 2020 | [68] |
| HeLa cells                 | CDYL KO              | 3734               | 14,311    | Negatively charged glutamic acid (E) at −1 and +1 positions | Mitochondria (18%), cytoplasm and nucleus (roughly equal) | Translation initiation, RNA splicing, and amino acid metabolism | RPA1, APEX1, and HP1 | 2020 | [52] |
| HCT116 cells              | P300 KO              | 392                | 816       | Glutamic acid (E) at −4, −1, +1, and +4 positions, valine (V) at +2 position | Nucleus (69%), cytosol (89%), and mitochondria (19%) | Co-translational protein targeting to membrane, nuclear-transcribed mRNA catabolic process, and translational initiation | LMNB2 | 2018 | [61] |
| PBMCs of patients with hemodialysis maintenance | PBMCs were obtained and compared with normal controls | 347                | 1109      | Aspartic acid (D) and glutamine (E) at −1 to +4 position; lysine (K) at −10 to −5 and +10 to +5 positions | Upregulated crotonylated proteins: cytoplasm (46%), nucleus (14%), mitochondria (12%), and extracellular region (11%). Downregulated proteins: cytoplasm (57%), extracellular region (11%), mitochondria (9%), and nucleus (8%) | Upregulated proteins enriched in cellular metabolic process and cellular response to outside and downregulated proteins localized in cellular functions such as migration and development | N/A | 2018 | [56] |
| Biological samples analyzed | Experiment condition | Identified proteins | Kcr sites Amino acids flanking the identified Kcr sites | Subcellular localization | Biological function | Validated proteins by WB | Year | References |
|----------------------------|------------------------|---------------------|----------------------------------------------------------|--------------------------|---------------------|------------------------|-------|------------|
| H1299 cells                | Without any treatment  | 1024 2696           | Glutamate (E) residues at the −1 and +1 positions       | Cytoplasm (40%), nucleus (27%), and mitochondria (13%) | 18% of crotonylated proteins are involved in the cell process, 14% are involved in metabolism, 13% are single biological proteins, and 11% are biomodulatory proteins | NPM1, FHL1, ACTN1, Integrin β1, Vinculin, ERK2, CDK7, GAPDH, and OTUB1 | 2017  | [17]       |
| HeLa cells                 | Treated with NaCr      | 453 558             | Alanine (A) residues at +3 or −6 positions              | Nuclear proteins (62.3%), nucleoplasmic localization proteins (9.4%), other cellular components including plasma membrane, cytoplasm, Golgi apparatus, and unclassified proteins (28.3%) | RNA processing, nucleic acid metabolism, chromosome organization, gene expression, DNA conformational change and packaging, chromatin organization and cell cycle | CBX3, CBX5, MTA2, Cul4B, and HDAC1 | 2017  | [49]       |
|                            | Without any treatment  | 70 N/A N/A           | N/A                                                      | N/A                      | Various molecular functions and biological processes closely related to DNA and RNA metabolism and cell cycle | HDAC1 | 2017  | [49]       |
| A549 cells                 | Treated with SAHA      | 2021 8475           | N/A                                                      | Cytoplasm (35.7%), nucleus (27.4%) and mitochondria (13.9%) | Gene expression, transcription, protein folding, and a variety of metabolic processes | N/A | 2017  | [21]       |
| Others                     |                        |                     |                                                          |                          |                     |                        |       |            |
| T. gondii                  | RH strain              | 3735 1,061          | Isoleucine (I) and lysine (K) occurred upstream; leucine (L), lysine (K) and phenylalanine (F) occurred downstream | Concentrated in and around the nucleus | Related to protein translation, compound metabolism, and biosynthesis | N/A | 2019  | [71]       |
|                            | ME49 strain            | 3396 984            |                                                          |                          |                     |                        |       |            |
| The tea plants             | NH4+ deficiency/resupply | 971 2288           | Alanine (A), glutamate (E), and lysine (K) residues at surrounding positions | Chloroplast (36.8%), cytoplasm (33.0%), and nucleus (13.7%) | Photosynthesis, carbon fixation, and amino acid metabolism | N/A | 2019  | [60]       |
Table 3 (continued)

| Biological samples analyzed | Experiment condition | Identified proteins | Kcr sites the identified Kcr sites | Amino acids flanking the identified Kcr sites | Subcellular localization | Biological function | Validated proteins by WB | Year | References |
|----------------------------|----------------------|--------------------|------------------------------------|-------------------------------------------|--------------------------|---------------------|------------------------|------|------------|
| Oryza sativa L. japonica   | Without any treatment | 690                | 1265                               | Aspartic acid (D), glutamine (E), isoleucine (I), lysine (K), leucine (L), arginine (R), and valine (V) at surrounding positions | Chloroplast (51%), cytoskeleton (1%), endoplasmic reticulum (1%), and extracellularly located (2%) | Photosynthesis | N/A | 2018       | [70] |
| Carica papaya L.           | Without any treatment | 2120               | 5995                               | Alanine (A), aspartic acid (D), and glutamine (E) at −5 to −1 and +1 to +5 positions, while lysine (K) and arginine (R) at −10 to −5 and +5 to +10 positions | Chloroplast (713 peptides), cytosol (691 peptides), nucleus (290 peptides), and mitochondria (138 peptides) | Biosynthesis of antibiotics, carbon metabolism, biosynthesis of amino acids, and glycolysis | N/A | 2018       | [69] |
| Zebrafish embryos         | Without any treatment | 218                | 557                               | Hydrophobic (L, I, V) and acidic (D and E) amino acids at surrounding positions | Cytosol (58%), mitochondria (11%), extracellular matrix (8%), and plasma membrane (6%) | Translation, metabolic process, and diverse regulation of skeletal muscle contraction | N/A | 2018       | [18] |
| PAT                       | Degradation in Rhoodontula mucilaginosa | 629                | 1691                               | Negatively charged residues from −10 to +10 | Mitochondria (89%), cytosol (32%), and nucleus (14%) | Metabolic process, cellular process, and single-organism process with upregulated or downregulated proteins | N/A | 2018       | [54] |
| Nicotiana tabacum         | Without any treatment | 637                | 2044                               | Aspartic acid (D) and glutamine (E) at surrounding positions | Chloroplast (37%), cytosol (30%), nucleus (12%), and mitochondria (5%) | Carbon metabolism, the citrate cycle, glycolysis, and photosynthesis the biosynthesis of amino acids | N/A | 2017       | [25] |

A549 human lung adenocarcinoma cells, CDYL chromodomain Y-like protein, HCT116 human colon cancer cells, HeLa human cervical adenocarcinoma cells, HT29 human lung adenocarcinoma cells, IgAN immunoglobulin A nephropathy, KO knockout, NaCr sodium crotonate N/A not available, PAT patulin, PBMCs peripheral blood mononuclear cells, SAHA suberoylanilide hydroxamic acid, T. gondii toxoplasma gondii parasites
changes of crotonylation of non-histone proteins in homologous recombination-mediated DNA repair [52].

Chemical labeling can be used in samples that cannot be metabolically labeled, and several samples can be quantified in parallel, for example, by tandem mass tags (TMTs) [54] or by isobaric tags for relative and absolute quantification (iTRAQ) [55, 56]. However, due to the influence of labeled groups, the identification flux was lower than the label-free approach.

Label-free quantification allows direct identification and quantification of proteins in a large scale, usually requiring analysis of a sample in triplicate to ensure that the measured differences are statistically significant [17, 20]. However, the accuracy of label-free quantification is slightly worse than that of the labeled quantification, because the former may be affected by the stability of mass spectrometry and other factors. In recent years, mass spectrometry techniques with increased stability and repeatability, as well as greatly improved computational algorithms for quantification of MS data, have made label-free quantification an attractive option [57–59].

To reduce the complexity of samples, after digestion, the trypsin polypeptides can be divided into several components by high-pH reversed-phase fractionation (RPF), known as sample fractionation. To increase the depth of the crotonylation analysis, immunoaffinity purification is usually used, in which the pan-Kcr antibody is immobilized to a resin bead and selectively bound to the crotonylated tryptic peptides and is then eluted [17, 52, 60, 61]. Enrichment of crotonylated peptides is usually combined with sample fractionation to improve the efficiency of the next mass spectrometry (MS). Finally, the enriched peptides are usually separated by liquid chromatography and ionized in the electrospray source, and entered into the mass spectrometer for analysis. High-resolution and high-quality precision analyzers can detect hundreds or thousands of different molecular features in a single LC–MS experiment, but only a small fraction of which can be identified and quantified [62]. The abundance of these eluted peptides, which range over many orders of magnitude, is a formidable analytical challenge that has been driving the progress of faster and more sensitive instruments and detection modes over the past decades [63–65]. For instance, a scan mode termed parallel accumulation-serial fragmentation (PASEF) has recently been demonstrated to increase sequencing speed exponentially without loss of sensitivity [66, 67]. All these technological advances are high performance additions to the technology toolbox in crotonylomics.

Other experiments can also be used to verify the results of crotonylomics, such as western blotting and immunofluorescent staining. Immunofluorescent staining shows that crotonylated proteins are widely located in the cytoplasm and nuclei of H1299 and HeLa cells [17]. In addition, crotonylated proteins are widely found in a variety of tissues of mice, including lung, kidney, liver, colon, uterus, and ovary. Although these methods are not as efficient as MS, they can be used to verify the conclusions of large-scale experiments, such as NPM1, FHL1, ACTN1, integrin β1, ERK2, and CDK1, which are considered to be crotonylated as assayed by MS and can also be detected by western blotting [17].

**Proteomic characteristics of crotonylation**

Due to the progress of technology, the field of lysine crotonylation has developed rapidly in the past few years. Crotonylation was initially found to occur mainly in histones. However, a significant conclusion from recent proteomic studies is that most crotonylation events occur on non-histone proteins [17, 52, 56, 61, 68]. MS-based proteomics methods are now used in a variety of organisms not limited to humans, resulting in the identification of thousands of new crotonylation sites (Table 3). These studies show that crotonylation sites are often conserved in different organisms [18, 60, 69–71], thus it is obvious that crotonylation can be regarded as a protein modification that exists prevalently in all fields of life. Crotonylated non-histone proteins are widely distributed in subcellular compartments and participate in a variety of important cellular functions, signal pathways, and variant biological activities (Table 3).

Motifs refer to some specific amino acids sequences which localize near the lysine acylation site and are generally highly conserved. The identification of the sequence of modification sites and the study of the corresponding model peptides provides clues for predicting the potential modification sites of new proteins. For instance, motif analysis is often used to predict potential kinase phosphorylation sites in bioinformatics [68, 72]. The amino acid sequences of motifs have been extracted from the upstream and downstream of the crotonylated lysine residue sites, which can describe the sequence commonness around the crotonylation sites. The studies given in Table 3 describe the amino acid sequence background of the crotonylation sites in eukaryotes. The highly conserved amino acids in these motifs, that is, E and D [17, 25, 52, 56], are negatively charged [54], which may represent the specific amino acid bias near the histone and non-histone crotonylation sites. Although specific amino acids are preferred near crotonylated lysine, no clear crotonylated amino acid sequence is found, which may be due to the diversity of KCT which targets these sites.

Quantitative analysis can also help to find the function-related crotonylation sites which are regulated under
specific conditions. Sodium crotonate (NACR) can induce protein crotonylation through enzymatic reaction. HDAC1, an endogenously crotonylated protein, can be identified in HeLa cells under conventional cell culture conditions. In addition to HDAC1, after NACR treatment, CBX3, CBX5, MTA2, and Cul4B are also crotonylated, while NACR does not affect the expression of these proteins [49]. In A549 cells treated with SAHA, the proteins with down-regulated Kcr levels were highly enriched in metabolism, while the proteins with up-regulated Kcr level were enriched in biological processes such as protein folding, RNA stability, gene expression, and transcription [21]. It can be anticipated that quantitative proteomics will continue to play an important role in deciphering the cellular role of crotonylation.

**Biological functions of non-histone protein crotonylation**

The widespread existence of non-histone protein crotonylation draws attention to the more general pathway control layer mediated by protein crotonylation. Positively charged lysine residues are often involved in protein–protein interaction and protein catalytic activity. Crotonylation neutralizes the positive charge of lysine, thus affecting many aspects of protein function, such as gene transcription, DNA damage response, enzymes regulation, metabolic pathways, cell cycle, and localization of heterochromatin in the cell. Different protein crotonylation could result in different consequences, depending on the functional nature of the proteins.

**Gene transcription**

Kcr is a specific marker of sex-linked genes [73]. Histone crotonylation was first found to be directly related to gene transcription [7]. Histone crotonylation specifically labels enhancers and transcriptional initiation sites of active genes in human somatic genomes and male mouse germ cell genomes, affects the structure of chromatin, and promotes the prolongation of histone substitution in sperm cells [7, 74, 75]. Although the knowledges about the functions of protein crotonylation are largely from studies of histone crotonylation, that of the non-histone crotonylation have been increasingly discovered. Following are two examples about the roles of non-histone crotonylation in gene transcription.

**P300/CBP**

As a typical lysine crotonyltransferase (KCT), P300-mediated crotonylation can promote transcription [31, 61]. In addition, crotonyl-CoA can stimulate transcription more effectively than acetyl-CoA. It has been found that CBP/P300 I1395G mutant (only having KCT activity but not KAT activity) can replace endogenous CBP/P300 KCT to crotonylate promoter and enhance intracellular transcriptional activation, thus improving TGF-β1-induced activation of PAI1 and SMAD7 [32]. Stimulation of macrophages by LPS initiates a transcriptional program that requires P300 recruitment to many sites of the downstream gene.

**P53**

P53 is a tumor suppressor protein which binds to specific DNA sequences and transcriptionally activates target genes to regulate several key cellular processes, including cell cycle control, apoptosis, and DNA repair in response to genotoxic stress [76]. Acetylation of P53 and its role in regulating tumor suppression have been widely explored [77]. The acetylation at multiple lysine residues of P53 by P300/CBP promotes p53 gene transcription and the expression of proapoptotic genes [78]. At the same time, acetylation of the lysine residues of P53 stabilizes P53 by unbinding P53 to its E3 ubiquitin ligase MDM2 [79], while MDM2 in turn recruits HDAC1 to deacetylate P53 and reduce its stability [80]. Large-scale identification of protein crotonylation reveals that P53 can also be crotonylated. However, whether crotonylation of P53 has a lysine residue similar to acetylation and how crotonylation of P53 affects its activity remain to be determined [49].

**DNA repair**

DNA, which carries all the instructions for organisms to survive and reproduce, must remain unharmed in order to replicate accurately before cell division. Therefore, when cells undergo thousands of different types of DNA lesions that endanger the stability of the genome, the lesions must be faithfully repaired with the help of DNA repair proteins. These DNA repair proteins are often mutated in human tumors [81]. In order to cope with a large number of endogenous and exogenous DNA lesions, cells have evolved intertwined but highly orchestrated DNA damage response (DDR) networks, including the signal cascade response that senses DNA lesions and activates downstream cellular pathways to repair damaged DNA [82]. Of note, histone and non-histone PTM play a central role in labeling damaged DNA and stimulating DNA lesion response proteins to recruit and accumulate to DNA breakpoints to promote DNA repair. In this aspect, crotonylation of proteins takes a key role for DNA repair.

As a single-stranded DNA (ssDNA)-binding protein in eukaryotic cells, the replicative protein A (RPA) plays a key role in DNA metabolism in meiosis, such as DNA replication, repair, and homologous recombination (HR). Human RPA is a heterotrimer composed of RPA1, RPA2, and RPA3. Among them, RPA1 is responsible for the binding and interaction of ssDNA and DNA
metabolism-related factors [83, 84]. RPA1 interacts with Mre11-Rad50-NBS1 (MRN) complex to promote ssDNA terminal excision during HR, and cooperates with RAD51, BRCA2, and RAD52 to stimulate strand exchange [85, 86]. In addition, RPA1 interacts with WRN and DNA2L to stimulate WRN-mediated double-stranded DNA terminal dissociation and DNA2L-mediated ssDNA degradation [87]. RPA1 is regulated by various PTMs such as acetylation, phosphorylation, ubiquitin, and SUMOylation during DNA metabolism [88–90].

Upon DNA insults induced by camptothecin (CPT), the Kcr level of RPA1 is upregulated significantly. CPT is a recognized drug that induces replication fork collapse, S-phase arrest, and HR in DSB repair of tumor cells. CPT can cancel the reconnection activity of topoisomerase I and produce double-strand break (DSB) [91]. CPT treatment results in a significant increase of Kcr levels in RPA1-K88 and RPA1K379, but only a slight increase in the Kcr level of RPA1-K595. Since both K379 and K595 are located in the DNA binding domain of RPA1, the K379cr and K595cr of RPA1 may be more important to the ssDNA binding ability of RPA1 [52].

The Kcr of RPA1 plays a key role in homologous recombination-mediated DNA repair. In the process of CPT-induced DSB, the Kcr of RPA1 promotes its ssDNA binding ability, recruits RPA1 at the site of CPT-induced DNA injury, enhances its interaction with HR factors (such as BLM, DNA2L, Mre11, NBS1, and RAD51), facilitates RAD51 foci formation upon CPT insult, and promotes the formation of ssDNA triggered by CPT by interacting with several components of the resection mechanism including MRN complex, BLM, and DNA2L [52]. In HeLa cells treated with CPT, downregulation of RPA1 gene causes a significant increase in apoptosis which can be rescued by the overexpression of wild-type RPA1 rather than mutation of the Kcr sites of RPA1 [52]. It is suggested that the Kcr of RPA1 possibly makes cells survive and resistant to apoptosis under the condition of DNA damage (Fig. 3).

CDYL may regulate RPA1 Kcr dynamics in response to CPT insult. CDYL is required for HR-mediated DNA repair by promoting homology-directed repair of DSBs using “traffic-light reporter” system [92] and coordinating with histone chaperone CAF-1 and DNA helicase minichromosome maintenance (MCM) for proper chromatin structure at damage sites [91]. Although studies have shown that Kcr of RPA1 at K88, K379, and K595 are all negatively regulated by CDYL [52], the possibility cannot be ruled out that RPA1 Kcr may be regulated by other factors, such as crotonyltransferase and/or decrotonylases, which may also contribute to the dynamic changes of RPA1 Kcr after CPT treatment.

**Enzyme regulation**

Large-scale identifications indicate that many enzymes are crotonylated in certain conditions and crotonylation may affect enzyme activities. For example, after treatment with a HDAC inhibitor SAHA in A549 cells, the crotonylation levels of 30 proteins are upregulated at 40 lysine sites, these proteins include aldo–keto reductase, alpha-enolase, acyl-CoA-binding protein, heat shock protein (HSP) 90-alpha, L-lactate dehydrogenase, ribosomal L1 domain-containing protein 1, splicing factor, and 40S ribosomal proteins [21]. However, the crotonylated lysine sites of other enzymes are downregulated, such as K280 on aldehyde dehydrogenase and K419 on very long-chain acyl-CoA synthetase [21]. The activity of a certain enzyme is increased by crotonylation, but not by upregulation of enzyme expression [49]. Expression of HA labeled HDAC1 in HeLa cells is not affected by NaCr or by combined treatment of HDAC inhibitor TSA and nicotinamide (NAM), but NaCr promotes the crotonylation of HDAC1, and the decrotonylase activity of crotonylated HDAC1 on its histone substrate is lower than that of unmodified
HDAC1. These studies reveal the importance of crotonylation in the regulation of enzyme activity.

Metabolic pathways
Enrichment analyses based on GO annotation, KEGG pathway, and Pfam domain analysis show that crotonylated proteins participate in a variety of important cellular pathways and perform different functions. Crotonylated proteins are involved in many metabolic pathways, including ribosome, spliceosome, proteasome, and Parkinson’s disease pathway, and are significantly enriched in ribosomal structure, translation factor activity, and adenyly nucleotide binding in the human lung adenocarcinoma cell line H1299 [49]. Bioinformatics analysis shows that a considerable number of crotonylated non-histone proteins in human specimens are involved in a wide range of metabolic processes, including DNA and RNA metabolism (Table 3). Crotonylated proteins associated with metabolic pathways have a wide range of significance among different species [69]. Crotonylation in papaya is involved in carbon metabolism, antibiotic biosynthesis, amino acid biosynthesis, and glycolysis. In particular, forty crotonylated enzymes have been identified in the amino acid metabolic pathways, suggesting that crotonylation has a potentially conserved function in the regulation of amino acid metabolism in papaya [69].

Cell cycle
Crotonylated proteins are enriched in the cell cycle protein network. As the components of histone deacetylase complexes, ubiquitin ligase RNF2 and binding enzymes UBE2E1, NCOR1, and RBPB4 are thought to inhibit histone acetylation and can be crotonylated in the regulation process of chromatin organization and cell cycle [49].

In metazoans, members of the MCM family 2–7 form a hexamer complex at the DNA replication fork, which determines the initiation of DNA replication [93]. After NaCr treatment, the MCM protein in chromatin decreases significantly, and MCM3 is crotonylated [49], which indicates that DNA replication may be affected by protein crotonylation. The combination of these factors may inhibit DNA replication and affect the cell cycle.

Localization of heterochromatin
Heterochromatin protein 1α (Hp1α), also known as chromobox homolog 5 (CBX5), is a member of the heterochromatin family and is enriched in heterochromatin via binding with methylated histones [94]. The crotonylation of Hp1α (CBX5) changes its redistribution in the nucleus and reduces the binding to methylated H3K9, the latter is highly enriched in heterochromatin [49]. It is well known that Hp1α acetylation also reduces its binding to methylated H3K9 [95], suggesting that crotonylation of some proteins share similar functional role as acetylation.

Non-histone protein crotonylation in disease and as potential therapeutic targets
Misregulation of non-histone protein crotonylation is associated with a variety of human diseases, thus crotonylation-related proteins are potential therapeutic targets. Targeted treatments with small inhibitors of HDACs, KATs, and bromine domain proteins have become attractive therapeutic strategies. Although the biological effects of HDACs, KATs, and bromine domain inhibitors are usually related to histone acetylation, these drugs also regulate crotonylation, thus it is possible that non-histone crotonylation is also involved in their cellular effects.

Cancer
Cancer-related proteins are affected by Kcr through CBP/P300. By comparing the global quantitative proteomics of p300 knockout cells and wild cells, it is found that the protein biomarkers of cancer in 4.5% of the EDRN database are crotonylated, and 32 Kcr proteins are related to cancer genes, accounting for 5.9% of the total genes of COSMIC cancer gene bank, and 6 target proteins of P300 are identified as cancer gene related proteins [61].

Pairing the tumor specimens with adjacent normal tissues, immunohistochemical staining shows that lysine crotonylation occurs in both cytoplasm and nucleus, and the level of lysine crotonylation is downregulated in liver cancer, gastric cancer, and renal carcinoma, but is upregulated in thyroid, esophagus, colon, pancreas, and lung cancers [96]. Interestingly, in hepatocellular carcinoma, lysine crotonylation level is associated with the stage of tumor lymph node metastasis (TNM).

Knocking down HDAC or adding HDAC inhibitor TSA can increase the level of crotonylation and inhibit the motility and proliferation of hepatocellular carcinoma cells [96], suggesting that HDAC inhibitors may target Kcr. HDAC has been developed to a potential target for oncology therapy. Extensive efforts over the past 20 years have produced dozens of HDAC inhibitors, of which four HDAC inhibitors (vo insignat, romidessin, panabinostat, and belinostat) have been clinically approved for the treatment of cutaneous and peripheral T-cell lymphomas as well as multiple myeloma, and has been tested in several other cancers [97].

As a well-known HDAC inhibitor, SAHA can inhibit HDACI, HDAC IIa, and HDAC IIb, and has been approved for the treatment of refractory cutaneous T-cell lymphoma (CTCL) [98]. The therapeutic effect of SAHA on other tumors, such as non-small cell lung cancer (NSCLC) and breast cancer, has also been confirmed.
In addition, a series of inhibitors of the unique \( \pi - \pi - \pi \) domain of readers on the Kcr recognition sites have been developed. For example, XL-13, a selective inhibitor of ENL YEATS, can interfere with ENL recruitment on chromatin when it binds to endogenous ENL and cooperatively downregulate oncogenes in acute leukemia with MLL rearrangement [102]. As mentioned above, SAHA can enhance the Kcr of histone and non-histone proteins, but the mechanism of Kcr in SAHA or reader in the treatment of cancer remains to be studied.

Major depressive disorder

BTBR T Lpr3tf/J (BTBR) mice have developmental disorders in the central nervous system and show many abnormal neuroanatomical structures. Compared with B6 mice, the global Kcr level of the cerebral cortex of BTBR mice is increased [103], suggesting that the functional relationship between crotonylation and brain diseases is of great significance. About 70 kDa of proteins in mouse brain extract can be recognized by pan Kcr antibodies, indicating the crotonylated non-histone proteins in the brain [104].

Crosstalk of lysine crotonylation with lysine acetylation

Herein crosstalk means that proteins are modified by multiple PTMs and these PTMs can interact with each other. PTM crosstalk can integrate different signals and greatly improve their regulation potential. Different PTM can lead to competitive crosstalk, fighting for the same lysine residue. Since Kcr and lysine acetylation (Kac) are catalyzed by the same enzymes and are removed by the same enzymes, we give a brief discussion on the crosstalk between Kcr and Kac below.

By comparing the data of crotonylation with that of acetylation on non-histone proteins in A549 cells, it is found that a total of 548 sites are simultaneously modified by both lysine crotonylation and acetylation. After SAHA treatment, 40 sites of 30 non-histone proteins are upregulated at the crotonylation and acetylation levels [21]. This result is consistent with the conclusions of other studies [36, 49], indicating that the two modifications can share a common set of proteins and sites. It is known that the acetyltransferase P300/CBP also catalyzes protein crotonylation, but how essentially the same transferase chooses between Kcr and Kac of the substrate warrants further study.

Whether histone lysine is crotonylated or acetylated depends on the relative concentration of intracellular crotonyl-CoA and acetyl-CoA [40]. In the case of nutritional exhaustion, the level of acetyl-CoA decreases significantly, while the proportion of crotonylated histones may increase to preserve the transcription of key genes during starvation. The fluctuation of protein crotonylation associated with metabolic status may be a normal function of cell physiology and, when destroyed, may lead to disease [105, 106]. Short-chain fatty acid metabolism occurs in intestinal microflora and affects the epigenetic characteristics of multiple proteins [107], including colonic histone Kcr which is dynamically regulated by the cell cycle [104]. When cell energy is exhausted, peroxisomal fatty acid \( \beta \)-oxidation and H3K9 crotonylation increase and is accompanied by a decrease in the levels of ATP and acetyl-CoA, as well as a decrease in the expression of ribosomal biogenic genes [108]. In this metabolic state, the crotonyl reader Taf14 YEATS domain inhibits the expression of growth-promoting genes to adapt to the low-energy state of cells [109]. We speculate that crotonylation of non-histone protein is also regulated by the relative concentrations of crotonyl-CoA and acetyl-CoA to adapt to the changes of cell energy environment.

Future perspectives

MS-based quantitative proteomics defines an initial reference framework for comprehensive Kcr studies and helps to our understanding on non-histone protein crotonylation. The latest developments in high-resolution nuclear magnetic resonance (NMR), genome engineering, novel bioinformatics tools, and selective chemical probe development provide exciting opportunities for system-wide and detailed mechanism studies to meet future challenges [110–114].

Although crotonylation and acetylation share certain enzymes, the properties of many newly discovered enzymes that regulate non-histone protein crotonylation are still unknown. Whether the enzymes catalyze in a site-selective manner or the enzymes shuttle in different cell compartments are interesting issues to be studied in the future.

Non-histone protein crotonylation does regulate many metabolic enzymes, but the dynamic regulation mechanism of the modification sites has not been deeply studied. Future work may find more evidence on how non-histone crotonylation affects protein and cell function. Crotonylation has been proved related with many human diseases, which may open an attractive field for medical intervention. We further expect that non-histone Kcr may be a contributor to disease development and a therapeutic target. Similar to other rapidly developing research fields, studies on non-histone protein crotonylation in the near future may face both challenges and exciting discoveries.

Abbreviations

ACSS2: Acyl-CoA synthetase short chain family member 2; AF9: ALL1 fused gene from chromosome 9; CBX5: Chromobox homolog 5; CDYL: Chromodomain Y-like; CoA: Coenzyme A; CPT: Camptothecin; CTCL: Cutaneous T-cell
lymphoma, DOR: DNA damage response; DDXS: DEAD-box helicase S; DPF: Double plant homeodomain finger; DSB: Double-strand break; GCNS: General control nonrepressed-protein S; GNAT: GCNS-related N-acetyltransferase; HDAC: Histone deacetylase; hMOF: Human males absent on the first; H1p1a: Heterochromatin protein 1a; HR: Homologous recombination; HSP: Heat shock protein; IgAN: Immunoglobulin A nephropathy; InsP: Inositol phosphates; IR: Ionizing radiation; Kac: Lysine acetylation; Kcr: Lysine crotonylation; KDCR: Lysine decrotonylase; LC-MS/MS: Liquid chromatography coupled to tandem MS; LTR: Long-terminal repeat; MCM: Minichromosome maintenance; MOZ: Monocytic leukemia zinc-finger protein; NAP: N-Acetylphosphatase; KCT: Lysine crotonyltransferase; MCT: Lysine malonyltransferase; MCM: Minichromosome maintenance; MOZ: Monocytic leukemia zinc-finger protein; mPFC: Medial prefrontal cortex; MS: Mass spectrometry; MYST: Moz Ybf2 Sas2 and Tip60; NaCr: Sodium cotromate; NMR: Nuclear magnetic resonance; NPM1: Nucleophosmin-1; NSCLC: Non-small cell lung cancer; PAT: Patulin; PBMC: Peripheral blood mononuclear cells; P300/CREB: P300/CREB-binding protein; PCAF: P300/CREB-associated factor; PHD: Plant homeodomain finger; PTM: Post-translational modification; RPA: Replicative protein A; SAHA: Suberoylanilide hydroxamic acid; SCFA: Short-chain fatty acid; SIRT: Sirtuin family deacetylase; TAF1: TATA binding protein-associated factor-1; TAF1: TATA binding protein-associated factor-1; T; gondii: Toxoplasma gondii parasites; TNM: Tumor, lymph node, metastasis; TSA: Trichostatin A.

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Authors’ contributions

JH: conception and writing of manuscript draft; JL: references preparation; LZ: DW: reading and revising manuscript; JC: revising and supervising submission. All authors read and approved the final manuscript.

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The authors declare that they have no competing interests in this work.

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