SUMOylation is a post-translational modification that regulates a multitude of cellular processes, including replication, cell-cycle progression, protein transport and the DNA damage response. Similar to ubiquitin, SUMO (small ubiquitin-like modifier) is covalently attached to target proteins in a reversible process via an enzymatic cascade. SUMOylation is essential for nearly all eukaryotic organisms, and deregulation of the SUMO system is associated with human diseases such as cancer and neurodegenerative diseases. Therefore, it is of great interest to understand the regulation and dynamics of this post-translational modification. Within the last decade, mass spectrometry analyses of SUMO proteomes have overcome several obstacles, greatly expanding the number of known SUMO target proteins. In this review, we briefly outline the basic concepts of the SUMO system, and discuss the potential of proteomic approaches to decipher SUMOylation patterns in order to understand the role of SUMO in health and disease.
cancer [10]. Deciphering the cellular SUMO network may therefore help us in further understanding the pathology of these diseases and in developing potent drugs. In this review, we briefly summarize the biochemistry of the SUMOylation machinery, and highlight the difficulties and great potential of proteomic approaches to uncover new SUMO targets, SUMOylation sites and cross-talk between post-translational modifications. The roles of SUMO during specific cellular processes have been discussed elsewhere [11–17].

The biochemistry of the SUMO system

SUMO proteins: similarities and differences

In contrast to yeast, which has a single SUMO protein called Suppressor of MIF2 mutations 3 (SMT3) [18], the human genome includes four SUMO genes, designated SUMO1, SUMO2, SUMO3 and SUMO4. While SUMO1 is distinct from the other family members (47% sequence identity between SUMO1 and SUMO2/3), mature SUMO2 and SUMO3 share a sequence identity of 97%, making them indistinguishable to currently available antibodies. Therefore, this sub-group is mostly referred to as SUMO2/3. All members of the SUMO family are expressed as precursor proteins, and first need to be processed by specific SUMO proteases, resulting in a free C-terminal di-glycine motif that is attached via an isopeptide bond to the ε-amino group of an internal lysine residue of the target protein. SUMO4, however, contains a specific proline residue (Pro90), preventing it from being processed by the known SUMO proteases [19]. It is currently unknown if and by what means mature SUMO4 is produced. The mature forms of SUMO1–3 are ubiquitously expressed. SUMO1 and SUMO2/3 have unique and overlapping target proteins [20,21]. In contrast to SUMO2/3, a major fraction of SUMO1 is conjugated to Ran GTPase-activating protein 1 under normal conditions, a mechanism that is essential for efficient nucleocytoplasmic transport [22,23]. Conversely, SUMO2/3 is mostly conjugated under cellular stress conditions such as heat shock [24]. Without stress, the cell contains a pool of unconjugated SUMO2/3, indicative of a cellular mechanism facilitating quick adaption of cells to stress arising from the extracellular environment. To further boost the SUMO signal during stress responses, the flexible N-terminal tail of SUMO2/3 contains a lysine residue (K11) situated within a SUMO consensus motif that is preferentially recognized by the SUMO conjugation machinery. This residue is the major acceptor site for SUMO2/3 chain formation [25]. SUMO1 lacks this regular consensus site, and therefore cannot efficiently form chains, but is instead considered as a chain terminator when incorporated in SUMO2/3 chains [26,27]. However, proteomic analyses have identified additional non-consensus acceptor sites within SUMO1 and SUMO2/3, suggesting a very complex chain formation pattern in cells [28,29]. To what extent these linkages are used in comparison with K11 linkages is currently not understood.

SUMO activating enzyme and SUMO conjugating enzyme

Similar to other ubiquitin-like modifiers, SUMO proteins are attached to target proteins via an enzymatic cascade (Fig. 1). In the first step, the C-terminus of the mature SUMO moiety is activated via ATP hydrolysis, resulting in a SUMO adenylate that is further attacked by the catalytic site of the SUMO activating enzyme dimer (SAE1/2, E1), forming a thioester with a cysteine residue in SAE2. In a second
step, the SUMO conjugating enzyme UBC9 (E2) binds to SAE2, and SUMO is transferred to its catalytic cysteine residue. In contrast to the ubiquitylation machinery, which comprises 35 conjugating enzymes with distinct functions and substrates, the SUMOylation cascade only contains UBC9 as a single E2. This highlights the important role of UBC9 as a key player in the SUMO signalling network. It is therefore not surprising that changes in the expression and activity of this enzyme have an extensive effect on regulation of the entire SUMO system, and must be fine-tuned via a multitude of additional signals. UBC9 has a preference for binding to proteins carrying a specific SUMO consensus motif (ΨKxE, where Ψ is a bulky hydrophobic amino acid) [30]. Consequently, in contrast to ubiquitin, bioinformatics tools for prediction of SUMO attachment sites in target proteins are available [31,32].

**SUMO ligases**

*In vitro* studies have shown that SUMO-loaded UBC9 alone is able to SUMOylate proteins bearing a SUMO consensus site [30]. However, under physiological conditions, this process is facilitated by SUMO ligases (E3), which confer specificity to the substrate and are able to promote SUMOylation of substrates even without the presence of a SUMOylation consensus motif. While the human genome includes approximately 600 genes encoding ubiquitin ligases [33], only very few SUMO ligases have been described so far. The first group to be discovered was the Siz/PIAS RING family, containing a characteristic zinc finger domain structurally related to the RING domain of ubiquitin ligases. Siz/PIAS SUMO ligases are involved in a multitude of cellular processes, such as the DNA damage response, cell-cycle control and transcriptional regulation [34,35]. However, their substrate specificity has been shown to be rather low [15]. Uncovering the control mechanisms regulating the expression, localization and activity of these ligases may provide insights into the larger picture of the SUMO response after certain stimuli such as heat shock and DNA damage, and is therefore currently an important area of research.

In contrast to the Siz/PIAS family, the unrelated large Ran-binding protein 2 specifically stabilizes the SUMOylated moiety of Ran GTPase-activating protein 1, and forms a highly stable complex with UBC9, which is essential for nucleoplasmic transport [36]. Ran-binding protein 2 promotes SUMOylation of other SUMO target proteins, such as SP100 and topoisomerase IIα, and therefore has many additional functions, specifically during mitosis [37,38].

Increasing numbers of proteins are thought to promote SUMO conjugation, such as the human polycomb protein Pc2/CBX4 [39], histone deacetylase 4 [40] and the tumor suppressor p14Arf [41]. These findings suggest that additional SUMO ligases remain to be uncovered, consistent with the current understanding of the ubiquitin system and the fact that many hundreds of SUMO target proteins have been identified so far.

Very recently, the Fanconi anaemia protein SLX4 was suggested to be a SUMO ligase that is essential for the response to global replication stress, and it was found to bind to UBC9 and SUMOylate both itself and the DNA repair factor XPF *in vitro* [42,43]. Interestingly, the functions of SLX4 are dependent on so-called SUMO interaction motifs (SIMs) present within the protein [43,44]. Such SIMs have been shown to promote SUMOylation of proteins even without the presence of a SUMO consensus site [45], and consist of large hydrophobic residues flanked by unstructured and exposed regions. Many proteins have already been shown to contain SIMs, and a recently developed bioinformatics tool facilitates the prediction of both SUMO consensus sites and SIMs in proteins of interest [32].

**SUMO proteases**

SUMO modification is a reversible process. Deconjugation of SUMO is catalysed by specific SUMO proteases. Only a small number of SUMO proteases have been identified so far, all of which are classified as cysteine isopeptidases. In mammals, a family of six SUMO-specific proteases (SENP1, 2, 3, 5, 6 and 7) are involved in the maturation and deconjugation of SUMO moieties [46,47]. While SENP5 has a preference for the SUMO3 precursor protein [48], SENP1 and SENP2 have been reported to process all SUMO isoforms to their mature forms with varying efficiency [46,49,50]. SENP1 has an additional unique feature, as it is mostly required for deconjugation of SUMO1 from substrates [51], whereas the other family members exhibit a strong preference for SUMO2/3 deconjugation, and SENP6 and SENP7 show particular specificity for disassembly of SUMO2/3 chains [52,53]. However, similar to SUMO ligases, the specificity of the SENPs for certain substrates is thought to be regulated in a spatial and temporal manner via regulation of protein amounts, localization and activity. Other SUMO proteases, such as Ubiquitin-Specific Peptidase-Like protein 1 [54], DeSumoylating Isopeptidase 1 and DeSumoylating Isopeptidase 2 [55], have
been identified but do not appear to be involved in changing global SUMOylation patterns.

**SUMO-targeted ubiquitin ligases**

As post-translational modifications represent a fast and reversible mechanism to alter the characteristics of a protein, it is logical to assume that these processes and the machinery required for modification are themselves subject to regulation via post-translational modifications. More than 500 post-translational modifications are known to date [56], demonstrating the immense potential of these signals in fine-tuning even minor biochemical processes within the cell. The picture becomes even more complicated when considering the plethora of post-translational modifications that work in concert to orchestrate various essential cellular processes.

The identification of a novel class of ubiquitin E3 ligases that specifically enhance ubiquitylation of SUMOylated proteins revealed an essential cellular mechanism involving cross-talk between two major post-translational modifications [57]. In humans, two such SUMO-targeted ubiquitin ligases have been identified, each one containing characteristic SIMs to specifically recognize SUMOylated proteins. Both SUMO-targeted ubiquitin ligases form SUMO–ubiquitin hybrid chains. RING finger protein 4 (RNF4), the smallest member of this enzyme class, functions as a homodimer and contains at least three SIMs, explaining its preference for target proteins modified by poly-SUMO chains comprising at least three SUMO moieties [58]. In addition to SUMO polymers, closely spaced mono-SUMOylation events may also form binding sites for SUMO-targeted ubiquitin ligases. Several substrates of RNF4 have been described to date, with promyelocytic leukemia protein (PML) and its oncogenic fusion variant PML–RARα being the most prominent [58,59]. PML is degraded in an arsenic trioxide-induced manner, resulting in disassembly of PML nuclear bodies. These subnuclear compartments contain many other SUMOylated proteins, such as Sp100 and Daxx, suggesting that these proteins are also subject to RNF4-mediated proteolysis [60–62].

The second and more recently identified human SUMO-targeted ubiquitin ligase is RING finger protein 111 (RNF111), or Arkadia, which contains at least three SIMs for recognition of poly-SUMO signals [63]. While RNF111 has also been implicated in PML degradation, it also catalyses very distinct non-proteolytic processes during the DNA damage response, where it has been shown to form K63-linked ubiquitin chains on SUMOylated Xeroderma Pigmentosum group C-complementing protein [64].

**Proteomic approaches to decipher the SUMO code**

**Difficulties and pitfalls in identifying SUMO target proteins**

Due to constant improvements in proteomic approaches, the number of known SUMO target proteins is greatly increasing, but still lags behind the number of target proteins found for some other post-translational modifications, such as phosphorylation and ubiquitylation. Several major difficulties impede efficient identification of SUMOylated proteins on a global scale. First, SUMO expression levels are much lower compared to ubiquitin, and the amount of SUMOylated target protein usually only represents a minor fraction of the entire pool of the protein. Therefore, SUMO targets must first be enriched via immunoprecipitation or pulldown experiments. Second, SUMO proteases are highly potent and must be inactivated in denaturing buffers containing SDS, guanidine hydrochloride or urea. However, the use of antibodies during enrichment requires partial refolding of proteins after lysis, potentially reactivating SUMO proteases and allowing them to remove some of the SUMO moieties. Third, identification of SUMOylation sites via mass spectrometry is highly challenging, due to the cumbersome C-terminal tryptic remnants of mammalian SUMO proteins that are as large as 32 amino acids for human and mouse SUMO2 and SUMO3, and 19 amino acids for human and mouse SUMO1, which are too large for efficient identification by mass spectrometry (Fig. 2E,F). Several strategies have been developed to circumvent these obstacles (Fig. 2), and each method has its clear advantages and disadvantages as detailed below.

**Identification of endogenous SUMO target proteins**

Almost 600 potential SUMO target proteins were identified under completely endogenous conditions, making use of monoclonal antibodies that specifically purify endogenous SUMO moieties [21]. The biggest advantage of this method is the possibility of applying it to a broad range of samples, such as primary cell lines, entire organs or rare patient material. Similarly, enrichment of SUMO target proteins using SIM traps may be used to identify endogenous SUMO targets, but is mostly limited to poly-SUMOylated and multi-SUMOylated proteins [28,65]. However, both methods are relatively costly, require large amounts of sample material, and the number of
SUMO targets identified under these conditions is considerably less compared to exogenous purification systems. Additionally, the endogenous methods are not suitable to efficiently identify SUMOylation sites, making it challenging to distinguish between real SUMO targets and fortuitously co-purifying proteins, although attempts to chemically shorten the tryptic remnant of endogenous SUMO2 have been described [66].

Use of tagged SUMO variants and the identification of SUMO sites

For more efficient SUMO target enrichment, the N-terminus of SUMO is most commonly fused to an epitope or affinity tag. This approach facilitates the purification of SUMOylated proteins using antibodies or affinity matrices that specifically bind to the tag. Subsequent digestion with either trypsin or the endoproteinase LysC, depending on the SUMO mutant used, results in shorter SUMO peptides to facilitate identification of SUMO sites via mass spectrometry.

**Fig. 2.** Proteomic approaches to identify SUMO targets. (A) Purification via SUMO-specific antibodies. Cells are lysed under denaturing conditions to inactivate SUMO proteases. Afterwards, samples are diluted to obtain mild buffer conditions and SUMOylated proteins are purified using SUMO-specific antibodies. (B) Purification via SIM traps. Cells are lysed in a mild buffer supplemented with iodoacetamide, and SUMOylated proteins are purified using the SIM-containing protein RNF4 as bait. (C) Purification with epitope tags. Cells expressing a tagged SUMO fusion protein are lysed in denaturing buffer. For subsequent immunoprecipitation of the SUMO target proteins using antibodies targeting the protein tag, samples are diluted to obtain mild buffer conditions. Finally, they are trypsinized and analysed via mass spectrometry. (D) Purification with affinity tags. Cells expressing SUMO tagged with affinity tags are lysed in denaturing buffer, and SUMO targets are purified using affinity matrices that specifically bind to the tag. Subsequently, proteins are trypsinized and analysed via mass spectrometry. (E) After trypsin digestion, the C-terminal fragments of mammalian SUMO family members are too large to efficiently map the SUMO-conjugated lysines in target proteins. To enable site-specific purification, protease cleavage sites are introduced in the C-termini of mammalian SUMO family members. SUMO target proteins modified with these mutant versions of SUMOs are fused to specific protein tags and purified as previously described for epitope or affinity tags. Subsequent digestion with either trypsin or the endoproteinase LysC, depending on the SUMO mutant employed, results in shorter SUMO peptides to facilitate identification of SUMO sites via mass spectrometry. (F) Alignment of the C-termini of mature SMT3 from yeast and mature human ubiquitin, SUMO1, SUMO2 and SUMO3, demonstrating the various lengths of the tryptic remnants remaining after cleavage. Arginine and lysine residues are highlighted in red. The mutations used to facilitate identification of SUMO2 sites are indicated by arrows.
samples, and has allowed us to identify more than 1600 SUMOylated proteins in human cells [29]. SUMO fusion proteins are either exogenously expressed from transgenes [69,73] or endogenously expressed using elegant knock-in approaches [8,72]. Therefore, these methods are restricted to specific cell types and organisms. Additionally, exogenous expression of SUMO fusion proteins may lead to higher SUMOylation levels of target proteins compared to endogenous conditions. Identification of SUMO target proteins via this method should ideally be confirmed using an endogenous approach. Several model organisms expressing tagged SUMO proteins have been developed, including yeast [69], Arabidopsis [73] and mouse [8]. Simultaneous application of tagged SUMO variants, together with quantification methods such as label-free quantification and stable isotope labeling with amino acids in cell culture, has uncovered major dynamics in the SUMOylation pattern [24,60,68,74].

Point mutations of these SUMO fusions, such as insertion of an additional cleavage site situated near the C-terminus (Q87R, T90R and T90K), result in shorter proteolytic fragments, which facilitate efficient mass spectrometric analysis. These SUMO mutants have been successfully used to identify SUMO attachment sites [68,72,75–78]. Introduction of additional mutations to specifically enrich for SUMOylated peptides in a two-step purification protocol allowed identification of over 4300 SUMOylation sites that dynamically change in response to various cellular treatments [29]. It has been confirmed that these mutations do not alter the overall conjugation efficiency of the SUMO proteins. However, it cannot be excluded that use of these SUMO mutant variants may alter SUMO conjugation to specific target proteins, particularly as one of the SUMO mutants used is deficient in forming SUMO chains. Therefore, it is recommended that SUMO sites are individually verified for each target protein before proceeding with further experiments. To circumvent the use of SUMO mutant variants, our laboratory has recently established a new site-specific SUMO proteomics methodology using His10-tagged wild-type SUMO and SENP2, leading to identification of more than 750 wild-type SUMOylation sites in HeLa cells [79]. Despite its great advantage in identifying wild-type SUMO2 sites, this method still utilizes an exogenously expressed SUMO fusion protein. The levels of exogenously expressed SUMO fusion proteins must be tightly controlled to prevent over-expression artefacts. To circumvent these pitfalls completely, development of similar methods using endogenous SUMO proteins to identify SUMO sites would be of considerable value, but this remains a major challenge.

The various site-specific analyses described above have allowed us to refine the canonical SUMO consensus motif and identify additional residues promoting SUMOylation. Several sites match the so-called inverted SUMO consensus motif or lie within a hydrophobic cluster SUMOylation motif bearing hydrophobic amino acids in close proximity to the modified lysine [29,75]. Interestingly, the percentage of sites situated within a SUMO consensus motif decreases after cellular stress, indicating that SUMO ligases that are active under these conditions may facilitate SUMOylation of non-consensus sites [29]. Alternatively, inactivation of SUMO proteases after cellular stress may lead to stabilization of SUMOylation of non-consensus sites.

![Fig. 3. Protein group modification via SUMO. In response to specific cellular or external stimuli, the activity and localization of the SUMO conjugation machinery is altered, leading to SUMOylation of target proteins with similar functions during the cellular response. This protein group modification triggers the formation of larger protein complexes via specific SUMO–SIM interactions. Increased activity of SUMO proteases reverses this process, leading to disassembly of these protein complexes.](image-url)
Proteomic analyses to unravel protein group modification

The proteomic analyses of SUMO target proteins performed to date clearly support the concept of protein group modification (Fig. 3). This model suggests that, after specific stimuli, an entire group of functionally related proteins is SUMOylated, allowing a quick and efficient cellular response \[14,80\]. Accordingly, treatment with DNA damaging agents such as methyl methanesulfonate and hydroxyurea induces SUMOylation of entire clusters of functionally related DNA repair proteins, chromatin modifiers and replication factors \[60,81,82\], while arresting cells in mitosis increases SUMOylation of a protein sub-group required for accurate chromosomal alignment and segregation \[68\]. This collective modification is mostly achieved via the presence of SUMO ligases and proteases that are differentially regulated in a spatial and temporal manner. Proteomic analyses of SUMO target proteins after depletion of these enzymes may provide additional insights into their substrate specificity and function in distinct cellular pathways. In further agreement with the concept of protein group modification, mass spectrometric analyses identified several protein complexes that contain two or more SUMOylated subunits, including chromatin remodeling complexes, histone deacetylases and spliceosomes \[29\]. It is believed

![Diagram](https://via.placeholder.com/150)

**Fig. 4.** Cross-talk between post-translational modifications as identified via mass spectrometry. (A) Several SUMO target proteins contain a so-called phosphorylation-dependent SUMO motif, in which the modified lysine residue is followed by a phosphorylated serine, usually five amino acids further downstream. A serine residue situated in a phosphorylation-dependent SUMO motif of the nuclear protein NOP58 is phosphorylated via casein kinase 2 (CK2), promoting UBC9 binding and subsequent SUMOylation of the indicated lysine residue. (B) Similar to phosphorylation, acetylation via specific acetyl transferases may induce SUMOylation of a protein, as described for histone H3. (C) Many lysine residues that were found as SUMO acceptor sites have also been shown to be ubiquitylated or acetylated, suggesting extensive competition between these modifiers. (D) Dozens of enzymes regulating post-translational protein modifications have been identified as SUMO target proteins in proteomic screens, including SUMO pathway components (S), or enzymes regulating other post-translational modifications, such as phosphorylation (P), ubiquitylation (Ub), methylation (Me) or acetylation (Ac).
that formation of many of these complexes is triggered and stabilized by the presence of SUMOylated sub-units and additional SIMs, representing a key mechanism to maintain genome stability and nuclear integrity [14,17,80].

Post-translational modification cross-talk identified by proteomic analyses

In addition to providing a comprehensive view of the SUMOylated proteome, proteomic approaches are a valuable tool in deciphering the vast network of post-translational modifications that work in concert to regulate specific cellular pathways (Fig. 4). Many components of the enzymatic machinery required to modify proteins are subject to SUMOylation, such as kinases, phosphotases, ubiquitin ligases and proteases, demethylases, and acetyltransferases [83]. Accordingly, the SUMO machinery itself has been found to be post-translationally regulated. SUMOylation and phosphorylation of UBC9 have both been shown to promote its enzymatic activity [45,84]. In contrast, acetylation of UBC9 specifically inhibits modification of substrates containing a so-called negatively charged SUMO consensus motif (ΨKxExxEEEE), providing a clear example of how post-translational modifications may confer substrate specificity to a relatively promiscuous enzyme [85]. Interestingly, several lysine residues of ubiquitin have been shown to be SUMOylated, indicative of the formation of hybrid chains consisting of the two post-translational modifications [29].

The possibility of identifying SUMO sites has enabled us to identify the strong dependency of SUMOylation events on other post-translational modifications. An obvious example of cross-talk between phosphorylation and SUMOylation is the existence of a phosphorylation-dependent SUMO consensus motif, in which a phosphorylated serine residue is generally situated five residues downstream of a lysine within a SUMO consensus motif [86]. Dozens of target proteins bearing such phosphorylation-dependent SUMO consensus motifs have been identified via mass spectrometry, and, for some of these proteins, both modifications were simultaneously present on the same peptide, indicative of a direct dependency. Indeed, mutagenesis experiments have shown that phosphorylation of these residues strongly increases SUMOylation of the relevant lysines [29,75]. Accordingly, an acetylation-dependent SUMOylation motif has recently been shown to be present in histones H3 and H4 [29]. Furthermore, many lysine residues identified as SUMOylated have also been shown to be acetylated or ubiquitylated, indicative of competitive usage of a subset of lysines by these post-translational modifications.

Conclusions and outlook

After 20 years of SUMO research, we are only beginning to grasp the enormous potential of this post-translational modification for regulating a vast number of cellular processes. Proteomic approaches have allowed us to identify hundreds of SUMO target proteins, but the number is still lagging behind findings for other essential post-translational modifications such as phosphorylation and ubiquitylation. However, the current proteomic analyses greatly help in understanding the network of SUMO targets within the cell, and further underline the concept of protein group modification. In addition, identification of specific SUMO sites and other post-translational modifications regulating SUMOylation events is facilitated by these mass spectrometric approaches. New purification methods that consecutively enrich SUMO and other post-translational modifications would further expand our knowledge about cross-talk between post-translational modifications, but are highly challenging and have not been successfully applied. Finally, it is of great importance to further improve purification methods to enrich endogenous SUMO proteins, to study relevant samples such as human patient material. Being able to identify SUMO target proteins, SUMO sites and components of the SUMO machinery that are deregulated in specific human diseases, such as cancer and neurodegenerative disorders, is an important step in developing new therapies and potent drugs.

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Author contributions

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