The HRASLS (PLA/AT) subfamily of enzymes

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Abstract

The H-RAS-like suppressor (HRASLS) subfamily consists of five enzymes (1–5) in humans and three (1, 3, and 5) in mice and rats that share sequence homology with lecithin:retinol acyltransferase (LRAT). All HRASLS family members possess in vitro phospholipid metabolizing abilities including phospholipase A₁/₂ (PLA₁/₂) activities and O-acyltransferase activities for the remodeling of glycerophospholipid acyl chains, as well as N-acyltransferase activities for the production of N-acylphosphatidylethanolamines. The in vivo biological activities of the HRASLS enzymes have not yet been fully investigated. Research to date indicates involvement of this subfamily in a wide array of biological processes and, as a consequence, these five enzymes have undergone extensive rediscovery and renaming within different fields of research. This review briefly describes the discovery of each of the HRASLS enzymes and their role in cancer, and discusses the biochemical function of each enzyme, as well as the biological role, if known. Gaps in current understanding are highlighted and suggestions for future research directions are discussed.

Keywords: H-RAS-like suppressor enzymes, Acyltransferase, Phospholipase A₁/₂, Class II tumour suppressor, N-acylphosphatidylethanolamine (NAPE), Pla2g16, TIG3, RARRES3, RIG1, iNAT

Introduction

The H-RAS-like suppressor (HRASLS) enzymes consist of five members (1–5) in humans and three (1, 3, and 5) in mice and rats [1, 2]. Also known as lecithin:retinol acyltransferase (LRAT)-like proteins due to their sequence homology to LRAT (Fig. 1) [3], the HRASLS enzymes comprise a vertebrate subfamily of the papain-like or NlpC/P60 thiol proteases that includes LRAT [4]. Enzymes in this superfamily have diverse functions, forming a conserved group of homologous proteins found broadly in the bacterial superkingdom, in DNA and positive strand RNA viruses, in archaea, and in eukaryotes [4].

HRASLS family members share several common elements (Figs. 1, 2 and 3), including a highly conserved NCEHFV sequence in the C-terminal region and similar basic structural motifs [2, 3]. Several residues in the NCEHFV sequence are critical for acylation and deacylation reactions, including the cysteine that acts as the active site nucleophile [2]. This residue is paired in HRASLS 2–5 with two N-terminal region histidines, and in HRASLS1 with N-terminal region histidine and asparagine residues, forming an unconventional catalytic triad that is characteristic of NlpC/P60 family members [4]. In this arrangement, the first histidine acts as a base to deprotonate the sulfhydryl group of the cysteine side chain, while the second polar residue acts to stabilize the correct orientation of the imidazole ring of the first histidine [2]. With the exception of HRASLS5, these proteins also contain a C-terminal transmembrane-spanning hydrophobic domain for endomembrane localization (Fig. 1) [5]. Currently, only the structures of HRASLS2, HRASLS3, and HRASLS4 have been reported. The crystal structures of HRASLS2 and HRASLS3 were solved by Golczak et al. in 2012. They were both shown to contain three α-helices well separated from a four-stranded antiparallel β-sheet, which resembles the basic structure of the NlpC/P60 thiol proteases that contain an α + β fold with segregated αβ segments (Fig. 2) [3, 4]. The NMR structure of HRASLS4 was solved by Wei et al. in 2015 and showed similar folding to that of HRASLS3, although the basic structure contained four α-helices and a six-stranded anti-parallel β-sheet (Fig. 2) [6].

Recent evidence has indicated a biochemical role for all HRASLS homologues in acyl chain remodeling of glycerophospholipids [3]. Using mass spectrometry analysis, a thioester intermediate has been detected to form
between cysteine 161 of purified LRAT and the acyl side-chain donated by phosphatidylcholine (PC) [7]. This suggests that the transient acylation of a reactive cysteine residue also occurs in the LRAT-like HRASLS enzymes to mediate reactions involving glycerophospholipids (Fig. 4) [3]. HRASLS1-5 all possess in vitro Ca^{2+} independent phospholipaseA_{1/2} (PLA_{1/2}) activities, which requires a pH of 8–9, dithiothreitol (DTT), and Nonidet P-40 for maximum function, and which are inhibited by iodoacetate [8–10]. The importance of DTT in particular suggests that preventing oxidation of sulfhydryl groups is critical for maintaining enzyme stability, which agrees with a predicted role for an active site cysteine in HRASLS enzymes [8]. Both PC and phosphatidylethanolamine (PE) act as substrates with PLA_{1} activity showing dominance over PLA_{2} activity for HRASLS1, 2, 4, and 5 [2, 8–11]. In regards to HRASLS3, there has been contradictory evidence of substrate specificity and positional preference for acyl hydrolysis, as well as other enzymatic characteristics [12–15]. Uyama et al. reported both PLA_{1/2} activities with PC and PE as substrates, but a preference for hydrolysis at the sn-1 position [15]. Duncan et al. also found that this enzyme shows both PLA_{1/2} activities with maximal phospholipase activity at pH 8, and they further identified a broad range of possible substrates, showing evidence of acyl hydrolase activity by HRASLS3/AdPLA against not only PC and PE, but also phosphatidylserine, and phosphatidylinositol, but not phosphatidic acid [14]. However,
in contrast to the findings of Uyama et al., Duncan et al. reported a preference for hydrolysis by this enzyme at the sn-2 position that is moderately stimulated by Ca\(^{2+}\) [14]. The inconsistencies within the literature have been proposed to be due to differences in the assay systems used [15]. Based on the evidence to date, it seems that this enzyme should best be considered a PLA\(^{1/2}\) [14–16].

The HRASLS family of proteins also possess O-acyltransferase activity, which describes the acyl-CoA independent transacylation of a lysophospholipid at either the sn-1 or sn-2 position using a fatty acyl chain donated by a phospholipid (Fig. 5) [1–3]. Each member of the HRASLS family has shown some preference for O-acylation of lysophosphatidylcholine (lyso PC) at the sn-1 position [1, 8, 10]. It has therefore been proposed that the HRASLS family be reclassified as a novel group of phospholipase A/acyltransferases (PLA/AT) 1–5 [1].

The HRASLS family of proteins also function in vitro to differing extents as N-acyltransferases for the production of N-acylphosphatidylethanolamines (NAPEs). HRASLS1-5 have been shown to catalyze the transfer
of an acyl chain from the \( sn-1 \) position of a glycerophospholipid to the amino group of PE producing NAPEs, which serve as precursors for N-acylethanolamines (NAEs) (Fig. 5) [1]. NAEs are a class of bioactive signaling molecules produced when a NAPE molecule is cleaved through hydrolysis, as catalyzed by NAPE-hydrolyzing phospholipase D (NAPE-PLD) (Fig. 5). NAEs have myriad biological roles, including modulation of inflammation, energy regulation, and nociception [1, 17]. Although little is yet known regarding the physiological significance of NAE regulation by HRASLS subfamily members, these enzymes are already known to play diverse biological roles in vivo. In particular, a function has been investigated for several members as class II tumour suppressor genes. Class II tumour suppressors are involved in the negative regulation of cell growth, and are often down-regulated in cancer cells secondary to other genetic or metabolic changes, but are not themselves mutated or deleted. As a result, they make attractive therapeutic targets, since they should be inducible if appropriate conditions or agents are identified [18–20]. Due to the study of HRASLS subfamily members in wide ranging fields, rampant renaming has occurred, with the result that there are now 27 different designations for these five enzymes (Table 1). Here, we identify and review the biochemical and biological role of these enzymes, and seek to disambiguate the current state of knowledge on this enzyme subfamily.

**Review**

**HRASLS1 (PLA/AT1)**

HRASLS1 was originally discovered by Akiyama et al. in 1999 through differential display between two mouse cell lines [21]. The 167-amino acid long protein was named A-C1, and was found to be 46 % homologous with a rat tumour suppressor protein called rat H-rev107 (now also called HRASLS3, among other names (see Table 1)) [11, 21]. The 168 aa-long human HRASLS1/A-C1 gene was also cloned, and was found to share 83 % identity with the mouse sequence [8, 22]. A tumour suppressor role for HRASLS1/A-C1 was tested directly and it was found that, indeed, overexpression of HRASLS1/A-C1 inhibited the growth of \( H\)-Ras-transformed NIH3T3 fibroblasts [21]. Additionally, expression of the human homolog was found to be reduced in gastric cancers from 41 different individuals due to hyper-methylation of CpG islands in the 5‘ region of the HRASLS1 gene, suggesting that it also plays a role as a tumour suppressor in human disease [23].

The enzymological role was first assessed over a decade after HRASLS1 was cloned [8]. Reports demonstrate a ratio of 1.6 [1] or higher [24] for human, mouse and rat HRASLS1/AC-1 PE N-acyltransferase activity to PLA\(_{1/2}\) activity, indicating greater specific N-acyltransferase activity in vitro [1, 24]. This is supported by metabolic labelling experiments in cells [1, 2, 8, 24].

The tissue distribution of HRASLS1 is distinguishable from other members of the HRASLS family of enzymes. In humans, mice, and rats, HRASLS1/A-C1/PLA/AT1 is
predominately expressed in testes, skeletal muscle, and heart [8]. Humans show relatively low expression of HRASLS1 in most other tissues [8], while mice and rats also show abundant expression in brain, with some evidence of expression in thymus, lung, stomach, kidney, and bone marrow cells [8, 21]. N-acylethanolamines (NAEs) are known to accumulate in response to acute injury in tissues, including the heart, brain, and testicles, suggesting a possible role for HRASLS-1 in mediating pathological events in these tissues [17]. Thus far, however, no studies have examined the physiological role of HRASLS1.

**HRASLS2 (PLA/AT2)**

HRASLS2 was first cloned from SW480 colon cancer cells [5]. This member of the HRASLS family is located
on chromosome 11 in humans, but is lacking from the rodent genome [5, 10]. The HRASLS2 gene transcript is detectable primarily in gastrointestinal tissues, including the colon, small intestine, and stomach, and is also expressed in the trachea and kidney [5]. Subcellular localization of HRASLS2 is primarily within the perinuclear region, where it shows a granular pattern [5], similar to that visualized for HRASLS1 [21].

A role for HRASLS2 in cancer and, specifically, in RAS-mediated cellular transformation, has been investigated. Transfection of HRASLS2 into HCT166 colon cancer cells, HeLa cervical cancer cells, and MCF-7...
breast cancer cells was shown to suppress colony formation [5]. As well, HRASLS2 over-expression suppressed the activation of wildtype and mutant RAS in HTA cervical cancer cells, resulting in increased cell death [5]. These effects were dependent on the C-terminal hydrophobic domain. Truncation of the last 26 amino acids containing this region eliminated the pro-apoptotic, anti-RAS, and growth suppressive activities of HRASLS2 [5]. In addition, loss of the putative C-terminal transmembrane domain also caused HRASLS2 to translocate, demonstrating increased intensity in the nucleus, which highlights the importance of this region in endomembrane localization [5].

The phospholipid metabolizing activities of HRASLS2 have been assessed using both cell homogenates and purified enzyme. HRASLS2 overexpression in COS-7 cells was found to increase in vitro PLA\(_{1/2}\) activity, with similar increases evident in assays conducted using either post-microsomal supernatants or microsomal pellets [10]. HRASLS2 was also shown to prefer to use fatty acids from the \(sn-1\) position of PC as acyl donors for its N-acyltransferase activities [1, 10]. In vitro, N-acyltransferase activity was found to be approximately 4-fold greater than PLA\(_{1/2}\) activity [1, 10]. Because of the preference for activity at the \(sn-1\) position, the major N-acyl species of NAPE produced by the overexpression of HRASLS2 in COS-7 cells tended to contain mostly saturated and monounsaturated acyl chains, with N-palmitoyl ethanolamide, N-stearoyl ethanolamide, and N-oleoyl ethanolamide forming the predominant NAEs generated, while N-arachidonoyl ethanolamide (i.e. anandamide), and its precursor N-arachidonoyl PE, showed only minimal elevation [1].

The physiological function of HRASLS2 is as-of-yet uncharacterized. Overexpression of HRASLS2 in clonal cell lines has been shown to decrease endogenous plasmenylethanolamine levels, which causes abnormal localization of peroxisomal proteins [1]. A similar result has also been observed for HRASLS3 [12]. Since the predominant activity of that enzyme is PLA\(_{1/2}\) rather than N-acyltransferase activity, it is unlikely that changes in NAPE/NAE levels or synthesis play a role in mediating peroxisomal events by HRASLS subfamily members. Further studies with \(Hrasls2\) gene ablation will be important to elucidate the biological role of this enzyme.

**HRASLS3 (PLA/AT3)**

Diverse studies on both the biological role and enzymatic function of HRASLS3 have led to the generation of a particularly large number of aliases, with at least ten different names recorded in the literature (Table 1) [9, 13, 25]. HRASLS3 was the first of the HRASLS subfamily of proteins to be discovered [11]. It was identified by subtractive hybridization performed between \(H\)-\(ras\) transformed rat fibroblasts and phenotypic revertants that had regained density-dependent growth inhibition, and thus was initially given the name \(H\)-\(ras\) revertant #107 (H-rev 107) to denote this [11]. In additional studies, HRASLS3/H-rev107 was categorized as a class II tumour suppressor [11, 26]. It has been suggested that HRASLS3 regulates oncogenic \(H\)-\(ras\)

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**Table 1** Aliases for the HRASLS family of enzymes

| HRASLS-1          | HRASLS-2          | HRASLS-3          | HRASLS-4          | HRASLS-5          |
|-------------------|-------------------|-------------------|-------------------|-------------------|
| **Aliases**       | PLA\(_{1/2}\)-2   | HRSL3             | HREV107           | HREV107-1         |
| A-C1              |                    | HRSL4             | TIG3              | RARRES3           |
| HRASLS            | PLA/AT-2          | HREV107           | RIG1              | HRLPS             |
| PLAT-1            | PLA/AT-1          | H-REV107-1        | iNAT              |                   |
| **Chromosomal location** | **Human 3q29** | **Human 11q12.3** | **Human 11q12.3** | **Human 11q13.2** |
| **NM accession No.** | **Mouse 16 B2** | **Mouse 19 A** | **Mouse 19 A** | **Mouse 19 A** |
| **GenBank**       | **Human NM_020386.4** | **Human NM_017878.1** | **Human NM_001128203.1** | **Human NM_004585.4** |
| **NP accession No.** | **Mouse NM_013751.5** | **Mouse NM_139269.2** | **Human NM_001324.1** | **Human NM_004585.4** |
| **GenBank**       | **Human NP_065119.2** | **Human NP_060348.1** | **Human NP_00121675.1** | **Human NP_004576.2** |
| **Mouse NP_038779.2** | **Mouse NP_644675.2** | **Mouse NP_080007.1** | **Mouse NP_080007.1** | **Mouse NP_080007.1** |

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through its ability to decrease cellular levels of ether-type lipids, such as plasmalogens and monoaalkyldiacylglycerols, by decreasing cellular levels of peroxisomes that are major sites for the synthesis of these lipids [12, 27]. Uyama et al. found that HRASLS3 binds to the peroxisomal chaperone protein Pex19p through its C-terminal and N-terminal hydrophobic domains [27]. Subsequently, Pex19p is inhibited from binding to peroxisomal membrane proteins such as Pex3p and Pex11p, which results in decreased peroxisomal biogenesis [27]. Additional studies, however, are still required to directly test the relative importance of peroxisomal biogenesis regulation in mediating effects of HRASLS3 on the H-ras-dependent growth of cancer cells.

HRASLS3 is downregulated in many cancers by methylation of the CpG-rich region in the promoter, which results in gene silencing [18, 28, 29]. In OVCAR-3 ovarian cancer cells, HRASLS3 has been shown to interact through its N-terminal proline-rich domain to ablate the catalytic activity of protein phosphatase 2A [25], causing caspase-9-dependent cell death. As well, HRASLS3 was found to inhibit cell migration and invasion when expressed in human NT2/D1 testicular cancer cells, and this function was suggested to relate to its ability to enhance the activity of prostaglandin D2 synthase [30]. At least one study has demonstrated, using chemical inhibitors, that the phospholipase activity of HRASLS3/H-rev107 is required for its anti-RAS effect [31]. This is also in agreement with an earlier finding that the growth-inhibitory and tumor-suppressing activity of HRASLS3/H-rev107 requires the C-terminal region [26] that is required for catalysis [16].

Not all studies, however, have found an inhibitory role for HRASLS3 in cancer. HRASLS3 does not inhibit, but instead stimulates, the proliferation of non-small cell lung carcinomas, contributing to tumour progression [29]. HRASLS3 has also been found to be a downstream target of mutant p53, with increased HRASLS3 levels reported in p53 mutant osteosarcomas [32]. As well, increased proliferation, migration, and invasion of osteosarcoma cells was reported following overexpression of HRASLS3 [32]. Additional studies on HRASLS3 active site mutants in transformed cell lines will help to isolate its lipid-enzymatic function from its function in tumour growth modulation.

The enzymatic activity of HRASLS3/RLP-3 was first investigated in 2007 by Jin et al., who demonstrated N-acyltransferase activity for this enzyme in vitro [9]. Subsequent work indicated the predominance of PLA over N- or O-acyltransferase activities for HRASLS3 in vitro [1, 8], and a detailed characterization of this enzyme as an adipose-specific phospholipase A2 (AdPLA) has been performed [14]. On the basis of studies performed using general phospholipase chemical inhibitors, it was suggested that this enzyme constituted the first member of an entirely new group of phospholipases A2, group XVI. As a result, the gene was subsequently renamed PLA2G16 [14].

Highly detailed analyses of human HRASLS3 have been performed in a series of elegant papers, including X-ray crystallographic analysis of its structure [3, 16, 33]. In the first structure analysis of the HRASLS family of proteins, Ren et al. demonstrated in 2010 that the phospholipase active site of HRASLS3 contains a Cys113-His25-His35 catalytic triad [33]. This finding was supported by Pang et al. in 2012 who showed that the phospholipase activity of HRASLS3 was mediated by this catalytic triad [16]. As is also seen in LRAT-NlpC/P60 family proteins, the active-site cysteine performs a nucleophile attack on the sn-1 and sn-2 acyl groups of phospholipids. In this reaction, the pK of Cys113 is too high to allow it to act as a nucleophile alone [16]. Thus, His25 assists in deprotonating the sulfhydryl side chain of Cys113, which effectively lowers the pK from 8.3-8.8 to 7.0, allowing it to function in the decylation of phospholipids. This study also provided mechanistic insight into the higher phospholipase activity relative to acyltransferase activity of HRASLS3 in particular. Evaluation of hydrogen/deuterium exchange at the active site of HRASLS3 suggested that during catalysis, water could readily access reactants, allowing the reactive intermediate to decompose to a free fatty acid and a lysophospholipid [16]. It remains to be determined whether varying degrees of hydrophobicity at the active site are related to the acyltransferase/phospholipase activity ratio of other HRASLS subfamily members. The C-terminal hydrophobic domain is critical for HRASLS3 activity. Truncation of the C-terminal hydrophobic domain results in loss of HRASLS3 from membranes, and a loss of HRASLS3-mediated phospholipase activity in vitro, indicating the critical nature of this hydrophobic span for interfacial catalysis [27].

The in vivo role of HRASLS3 has been studied in some depth [13]. HRASLS3/AdPLA is expressed predominantly in white adipose tissue (WAT), and also to a lesser extent in brown adipose tissue [13, 14], where it is a major regulator of lipolysis and is crucial for the development of obesity, as seen in an Hrasls3−/−/AdPLA−/− mouse model [13]. The ablation of Hrasls3/AdPLA causes a reduction in adipose tissue mass and triacylglycerol content, despite normal adipogenesis, as a result of constitutively elevated rates of lipolysis. The underlying molecular mechanism involves the regulation of lipolysis through the modulation of prostaglandin E2 (PGE2) levels and signaling. HRASLS3/AdPLA is responsible for 80% of adipocyte phospholipase activity, which is a major source of arachidonic acid that is used in the synthesis of eicosanoids [13]. As a result, loss of
HRASLS3/AdPLA results in a dramatic fall in levels of PGE$_2$ [13]. The primary receptor for PGE$_2$ in adipocytes is prostaglandin E receptor 3 (EP3), which is unique among EPs in that it is G$_{i/o}$-coupled, and therefore exerts a repressive action on adenyl cyclase in adipocytes when activated [34]. Thus, loss of HRASLS3/AdPLA in mice causes a decrease in PGE$_2$ levels and concomitant reduced activation of inhibitory EP3, with the result that generation of cAMP, activation of the hormone-sensitive lipase, and stimulation of lipolysis proceeds unchecked, preventing the accumulation of triacylglycerol and the development of genetic or diet-induced obesity [13]. Since HRASLS3/AdPLA is normally stimulated by insulin, increased activity by this enzyme may help to modulate the antilipolytic effects of insulin in adipocytes in the fed state [13]. The role of HRASLS3/AdPLA in adipocyte triglyceride metabolism suggests that inhibition may be a useful strategy to prevent and treat obesity [13, 16], and future studies will therefore likely focus on the design of inhibitors [16] and on strategies to regulate this enzyme.

HRASLS4 (PLA/AT4)

HRASLS4 was identified by three separate groups in 1998 [35], 2000 [36], and 2001 [37] as a retinoid-inducible anti-proliferative/class II tumour suppressor gene that goes by the alternate names Tazarotene-induced protein 3 (TIG3), Retinoic acid receptor responder protein 3 (RARRES3), and Retinoid-inducible gene 1 (RIG1) (see Table 1 for all aliases). HRASLS4 is homologous with HRASLS3/H-rev107 [35, 36] and, like that enzyme, has reduced expression in a wide variety of primary human tumours, including lymphoma, ureter, kidney, rectal, and uterine, and cancer cell lines, including HL-60 promyelocytic leukemia cells, HeLa cells, K-562 chronic myelogenous leukemia cells, SW480 colon carcinoma, A549 colon carcinoma, and G361 melanoma cells [35]. HRASLS4 also shows growth inhibitory effects when overexpressed in T47D Chinese hamster ovary cells [35]. A role for HRASLS4 in malignant disease continues to be studied extensively, and a comprehensive overview of this area is beyond the scope of the current review. However, it is of interest to mention some of the more recent findings. The HRASLS4 promoter contains a response element for the p53 tumour suppressor that is activated by wildtype but not mutant p53 [38]. Whether this feature exists for all HRASLS subfamily members has not yet been studied, but it would help to explain the almost universal down-regulation of these proteins in transformed cells.

Like other HRASLS subfamily members, HRASLS4 also inhibits H-RAS-mediated signalling [5, 39, 40]. The anti-cancer activity of HRASLS4 has been localized specifically to its action within the Golgi. While the C-terminal hydrophobic domain of HRASLS4 anchors this enzyme within both the endoplasmic reticulum and the Golgi apparatus, only Golgi-targeted HRASLS4 induces apoptosis in cancer cells [39]. The phospholipid metabolizing activity of HRASLS4 appears to be important for its anti-cancer effects, particularly with regards to metastasis and invasion [41]. Similar to HRASLS3, HRASLS4 functionally interacts with prostaglandin D$_2$ synthase to augment the production of prostaglandin D$_2$ [42]. This function was found to be dependent on an intact C-terminal hydrophobic domain, and although not tested directly, it seems highly likely to be related to the phospholipase activity of this enzyme [42].

A role for HRASLS4 in phospholipid metabolism has been investigated [1, 10]. HRASLS4 functions in vitro as a Ca$_{2+}$ independent PLA$_{2}$/ [1, 8, 10]. Although N-acyltransferase activity in vitro has been shown to be minor or absent for this enzyme [10], HRASLS4 does significantly increase the cellular content of both NAPE and NAE in metabolic labelling experiments [1]. The functional significance of this activity with regards to the anti-cancer activity of this enzyme remains to be determined.

A physiological role for HRASLS4 has been investigated [43]. HRASLS4 is found in the suprabasal epidermis of skin, where it interacts with and activates another enzyme, transglutaminase I (TG1), which functions during terminal differentiation to form covalent bonds between proteins at the inner surface of the plasma membrane [44]. The action of TG1 is critical for production of the cornified envelope that maintains the epidermal barrier [44], and adenoviral delivery of HRASLS4 to cultured keratinocytes results in the activation of TG1, resulting in differentiation-like cell death, as well as the generation of cornified envelope-like structures [45, 46]. Reduced expression of HRASLS4 is evident in hyperproliferative conditions such as psoriasis [47]. An additional role for HRASLS4 in keratinocyte differentiation is suggested by findings that this protein also localizes to the centrosome, where it affects microtubule kinetics and cell division [43, 48]. Interaction between the centromere and HRASLS4 has been found to involve a central region of the enzyme that contains the NCEHFV sequence [43]. However, whether the phospholipid metabolizing abilities of this enzyme, per se, are critical for its various physiological roles in skin cells, remains to be clearly characterized.

HRASLS5 (PLA/AT5)

HRASLS5 was first investigated as part of a search for new Ca$_{2+}$-independent enzymes active in the N-acylation of PE [9] (also called RLP-1 and iNAT; see Table 1 for all aliases). HRASLS5 displays dominate N-acyltransferase
activity over both PLA₁₋₂ and O-acylation activities in vitro, and overexpression of HRASLS5 enhances formation of NAPE and NAE in cultured cells [1, 2, 8, 9]. However, HRASLS5 differs from other HRASLS family members in that it is predominantly present in the cytosol, likely due to the absence of the C-terminal hydrophobic span that is characteristic of HRASLS enzymes 1–4 (Figs. 1, 2, 3 and 4) [1, 9]. HRASLS5 also does not show a clear preference for abstraction of acyl groups from either the sn-1 or sn-2 positions of PC during N-acylation reactions [9]. It has been suggested that HRASLS5 may be involved in the production of anandamide (arachidonoyl ethanolamide) because of its capacity to utilize fatty acyl chains from the sn-1 or sn-2 position of phospholipids [9]. However, studies have yet to characterize the nature of endogenous NAPE and NAE species produced by HRASLS5. Likewise, little is known regarding the physiological function of this enzyme. It has been identified in spermatocytes of developing rat testes, but its function in that tissue is currently unknown [49]. Similarly, studies have yet to investigate whether it has a role in cancer like the other HRASLS family members.

Conclusions

The five members of the HRASLS family of enzymes have been shown to have phospholipase A₁₋₂ activity, and O- and N-acyltransferase activity. In addition, studies generally report reduced expression of these enzymes in cancer cells, and demonstrate a direct anti-cancer role, although not all studies agree. Evidence from studies using mutated or truncated forms of HRASLS subfamily members suggests that the phospholipid metabolizing functions are required for some, but likely not all, of the effects observed in tumour cells. Further work should integrate recent advances in understanding the biochemical function of these enzymes to better understand mechanisms related to aberrant cell growth in neoplasia. Further studies should also focus on understanding the major physiological function of each homologue in cells and tissues, using recent advances in gene editing techniques as well as the generation of gene knockout mice. Human monozygotic twins with Poland Syndrome have been found to be heterozygous for a gene deletion event on chromosome 11 that removes HRASLS2-5, strongly suggesting a role for one or more of these genes in causation of this disorder [50]. Poland Syndrome is characterized by hypoplasia/aplasia of the pectoralis major muscle and other variable anomalies including hypoplasia/aplasia of mammary tissue and ribs, limited subcutaneous fat, and sternal anomalies [50]. Generation of gene knockout mice will be required to understand any putative role for this enzyme subfamily in normal physiology and in human disease.

Abbreviations

AdPLA: adipose-specific phospholipase A₂; DTT: dithiothreitol; HRASLS: H-RAS-like suppressor; LRAT: lecithin:retinol acyltransferase; lyso-PC: lysophosphatidylcholine; NAPE: N-acetylphosphatidylethanolamine; NAPE-PLD: N-acetylphosphatidylethanolamine-hydrolyzing phospholipase D; NAE: N-acylphosphatidylethanolamine; PLA₂/AT: phospholipases A₂/acyltransferases; PC: phosphatidylcholine; PE: phosphatidylethanolamine; PLA: phospholipase A.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

EBM reviewed the literature, carried out the sequence alignments, and drafted the manuscript. RMB created Figs. 4 and 5 and critically revised the text of the manuscript. RED was involved in the conception and design of the manuscript, and critically revised all elements. All authors read and approved the final manuscript.

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