Clusterin: a key player in cancer chemoresistance and its inhibition

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Abstract: Clusterin is a heterodimeric disulfide-linked glycoprotein (449 amino acids) isolated in the rat prostate after castration. It is widely distributed in different tissues and highly conserved in species. There are two isoforms (1 and 2) with antagonistic actions regarding apoptosis. Clusterin is implicated in a number of biological processes, including lipid transport, membrane recycling, cell adhesion, programmed cell death, and complement cascade, representing a truly multifunctional protein. Isoform 2 is overexpressed under cellular stress conditions and protects cells from apoptosis by impeding Bax actions on the mitochondrial membrane and exerts other protumor activities, like phosphatidylinositol 3-kinase/protein kinase B pathway activation, modulation of extracellular signal-regulated kinase 1/2 signaling and matrix metallopeptidase-9 expression, increased angiogenesis, modulation of the nuclear factor kappa B pathway, among others. Its overexpression should be considered as a nonspecific cellular response to a wide variety of tissue insults like cytotoxic chemotherapy, radiation, excess of free oxygen radicals, androgen or estrogen deprivation, etc. A review of the recent literature strongly suggests potential roles for custirsen in particular, and proapoptosis treatments in general, as novel modalities in cancer management. Inhibition of clusterin is known to increase the cytotoxic effects of chemotherapeutic agents, and custirsen, a second-generation antisense oligonucleotide that blocks clusterin, is being tested in a Phase III clinical trial after successful results were achieved in Phase II studies. A major issue in cancer evolution that remains unanswered is whether clusterin represents a driving force of tumorigenesis or a late phenomenon after chemotherapy. This review presents preclinical data that encourages trials in various types of cancer other than advanced castration-resistance prostate cancer and discusses briefly the appropriate timing for clusterin inhibition in the clinical context.

Keywords: custirsen, apoptosis, renal cell carcinoma, triple negative breast cancer, castration-resistance prostate cancer

Introduction
Castration in rats leads to involution of the ventral prostate, because this organ requires adequate concentration of androgens. One of the mechanisms involved in prostate involution is cell death. Castration elicits the appearance of several proteins in the prostate whose functions were unknown at the time. In 1989, a messenger (m)-RNA coding for sulfated glycoprotein-2 was identified in the prostate of castrated rats, peaking between days 3–4 after castration.1

Up to 1995, there was the concept that sulfated glycoprotein-2 (also dubbed apolipoprotein J or testosterone-repressed message-2), best known now as clusterin (CLU), was a protein associated with reduction in programmed cell death (apoptosis) in the prostate, but its exact role remained uncertain. This role was clarified by pioneering
studies performed by Sensibar et al working on LNCap cells, which are androgen-sensitive human prostatic cancer cells obtained from a human lymph node metastasis.

This research showed that CLU was associated with protection against induced cell death, reducing the cytotoxic effect of tumor necrosis factor (TNF-α) in LNCap cells. The overexpression of CLU, resulting in resistance to TNF-α-induced apoptosis, was not described prior to this research, which included the finding that there was a correlation between the expressed CLU and degree of resistance to TNF-α-induced apoptosis. The authors proposed the hypothesis that CLU overexpression can prevent cell death, whereas its depletion is responsible for the onset of cell death.

These data were later confirmed and amplified in a study showing that CLU knockdown in human cancer cells induced significant reduction of cellular growth and higher rates of spontaneous endogenous apoptosis.

**CLU gene**

The **CLU** gene is a well conserved gene during evolution, located in humans in chromosome 8 (8p21-p12) and it is expressed during tissue regression in many organs. Its overexpression should be considered as a nonspecific cellular response to a wide variety of tissue insults.

**CLU** is a single nine-exon gene expressing three protein forms with different subcellular localizations and diverse biological functions.

**Clusterin isoforms and structure**

However, things are not that simple regarding CLU, because we now know that the **CLU** gene has nine exons and there are at least three isoforms of clusterin: isoform 1, isoform 2, and isoform 11036. Exon 1 has three portions: 1a, 1c, and 1b.

Isoform 1 (nCLU, nuclear clusterin) is the product of the nine exons but contains only part 1a of exon 1, whereas isoform 2 (sCLU, secretory clusterin), contains the product of part 1b of exon 1 and isoform 11036 (cCLU, cytoplasmic CLU) contains part 1c of exon 1 (Figure 1). sCLU and nCLU are originated by the **CLU** gene through alternative splicing. nCLU is a slightly longer mRNA and with a completely different function: binding to Ku70 and promoting cell death by freeing Bax.

In summary, the **CLU** gene produces through alternative splicing three different proteins, two of which are sCLU, which prevents cell death, and nCLU, which promotes cell death. One CLU form is secreted (sCLU), whereas the other forms have a cytoplasmic (cCLU) or nuclear (nCLU) localization.

A unified nomenclature has been proposed. sCLU is reserved for the main gene transcript identified from the **CLU** gene locus. It has a predicted molecular mass of 50.1 kDa, which is detected as a 60 kDa glycosylated presecretory form of sCLU and is referred to as psCLU. psCLU is translated from the mRNA, including the nine gene exons. Once it forms the mature, secreted heterodimeric 75–80 kDa protein, it should be referred to as sCLU. Thus, mature sCLU is a heterodimeric disulfide-linked

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**Figure 1** Structure of **CLU** gene and transcription products. (A) **CLU** gene maps on chromosome 8. The position of exons and introns on the gene is shown. Black bands: exons, white bands: introns. (B) Exonic structure of **CLU** mRNA variants. Dark grey exon, exon 1a of isoform 1; medium grey exon, exon 1b of isoform 2; light grey exon, exon 1c of isoform 11036.


**Abbreviation:** CLU, clusterin.
Clusterin actions

CLU protects cells from various stressors that may induce apoptosis, like cytotoxic chemotherapy and androgen or estrogen deprivation. The cytoprotective role of CLU can be explained by its chaperone-like activity.12 CLU expression is low in most normal cells and is strongly stimulated by various stressors.5,13

How does CLU prevent cell death? The answer came in 2009: sCLU stabilizes the union between Ku70 and the apoptotic protein Bax, thereby preventing Bax from joining the mitochondrial outer membrane and exerting its proapoptotic activity, so that elevated sCLU levels may enhance tumorigenesis by interfering with Bax proapoptotic activities (Figure 2).14

Depletion of sCLU triggers two types of action: 1) p53 dependent; and 2) p53 independent (Figure 3).14-18 sCLU depletion activates p53 and alters the ratio of proapoptotic to antiapoptotic Bcl-2 protein family members, triggering mitochondrial dysfunction and apoptosis. However, Bcl-2 suppression is a p53-independent effect, such that knockdown of p53 only minimally suppressed cell death, but a pan-caspase inhibitor completely avoided cell death. Another mechanism involved in sCLU prosurvival activity is the upregulation of the phosphatidylinositol 3-kinase (PI3K)/protein kinase B (AKT) pathway15 and insulin-like growth factor (IGF)-1 activates the PI3K/AKT pathway through upregulation of sCLU.16

Besides its important role in apoptosis, sCLU has other actions related to tumor behavior. For example, in clear cell renal carcinoma (CCRC) sCLU regulates aggressive behavior by modulating extracellular signal-regulated kinase (ERK)1/2 signaling and matrix metallopeptidase (MMP)-9 expression.17 Knockdown of sCLU in cells showed reduced migration and invasion in vitro, as well as decreased metastatic potential in experimental metastasis.

Overexpression of CLU in epithelial ovarian cancer appears to be correlated with increased tumor angiogenesis, consistent with the established role of CLU as an oncogene in the biology of ovarian cancer.18 Also, sCLU increased chemoresistance in ovarian cancer cells.19

Furthermore, CLU has nuclear factor kappa B (NF-κB) regulatory actions.20 In prostate cancer cells, Zoubeidi et al.11 discovered sCLU’s ability to regulate ubiquitination and degradation rates of COMM domain-containing protein 1 and inhibitor of kappa B to enhance NF-κB nuclear translocation and transcriptional activity. Accordingly, sCLU knockdown stabilizes COMM domain-containing protein 1 and inhibitor of kappa B, sequesters NF-κB in the cytoplasm, and decreases transcription of many NF-κB regulated genes.

CLU is a protein widely distributed among cells, but certain situations like cellular stress, like chemotherapy, radiotherapy, or androgen depletion in prostate trigger CLU overexpression. CLU expression is under genetic and epigenetic control, the latter via methylation of its promoter.22

Many strategies used to kill cancer cells induce stress responses that activate survival pathways, including overexpression of sCLU, to promote emergence of a treatment resistant phenotype.

There is a strong association between CLU and glucose-regulated protein, according to Wang et al.23 Apparently CLU protects cells from endoplasmic reticulum stress-induced apoptosis, at least partially through interacting with glucose-regulated protein.

sCLU is a stress-activated cytoprotective chaperone upregulated by many varied anticancer therapies to confer treatment resistance when overexpressed.24 Using size exclusion chromatography and fluorescently labeled paclitaxel, Park et al25 demonstrated that clusterin binds to paclitaxel. These authors also showed that high levels of clusterin expression increased paclitaxel resistance in ovarian cancer cells by physically binding to paclitaxel, which may prevent paclitaxel from interacting with microtubules to induce apoptosis.
CLU expression is downregulated in human testicular seminoma, which is a highly sensitive tumor to radiotherapy and chemotherapy. The authors postulate that CLU expression in human testicular seminoma differs from CLU expression in other tumors and think that this fact may be the cause of the treatable characteristics of testicular seminoma.

There is no doubt nowadays that CLU has an important role in tumorigenesis and progression of human cancers, therefore it is a valid target for new cancer treatment modalities. Targeting the apoptosis pathway is a new and promising therapy.

Clusterin regulation
sCLU is overexpressed when stressor conditions are present in the cell, but at least in the case of ionizing radiation as a stressor condition, a regulatory pathway for CLU overexpression was identified: activation of ataxia telangiectasia-mutated kinase by endogenous or exogenous forms of DNA damage was required to relieve basal repression of IGF-1 transcription by the p53/NF-YA complex, leading to sCLU expression. Cells exposed to DNA-damaging agents had elevated IGF-1 expression, resulting in activation of IGF-1R signaling and sCLU induction. Overexpression of H-Ras caused repression of CLU expression. These authors support the concept that apoptotic processes, not oncogenic mutations, are responsible for increased clusterin expression in tumors.

p53 exerts a role in the transcriptional repression of sCLU. Loss of functional p53 in HCT116: p53(−/−) cells augmented CLU de novo synthesis after IR exposure. Repression of sCLU protein levels by p53 may be important for the cascade of p53-mediated events leading to cell death after IR or other cytotoxic agent exposure.

Clusterin inhibition
Custirsen (OGX-011; OncoGeneX Pharmaceuticals, Inc., Bothell, WA, USA) is a second-generation antisense oligonucleotide designed to block mRNA sCLU, producing depletion of the protein sCLU and improving clinical results in advanced prostatic cancer resistant to castration. Custirsen is currently in Phase III clinical trials. Even so, there is evidence that custirsen could be helpful in other kinds of tumors besides prostate cancer, such as CCRC, breast cancer, lung cancer, and cancer of the ovary, to mention a few.

The reasons for custirsen development
Custirsen binds to exon 1 of mRNA sCLU, blocking its translation. The development of this novel drug was due to findings like:

1. Renal carcinoma cells overexpressing sCLU became more resistant to apoptosis.

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**Figure 3** Mechanisms of clusterin action.
Notes: On the left side, two antiapoptotic mechanisms are described (p53-dependent and p53-independent). On the right side, other protumor mechanisms are not directly related with apoptosis.
Abbreviations: ERK, extracellular signal-regulated kinase; MMP, matrix metallopeptidase; NF, nuclear factor; PI3K, phosphatidylinositol 3-kinase; VEGF, vascular endothelial growth factor.
2. Oligonucleotides that target sCLU increase sensitivity to radiation in prostate cancer cells.42
3. Oligonucleotides that target sCLU increase sensitivity to chemotherapeutic agents.43

Possible custirsen activity in various tumors
Custirsen has been found to increase cytotoxicity when associated with docetaxel in prostate carcinoma cells,31 with sorafenib in CCRC,33 with cisplatin in CCRC,44 with gemcitabine and platinum compounds in lung cancer cells,30 and with paclitaxel.44

CLU and prostate cancer
It has been extensively investigated and there are many publications and reviews on this subject. Custirsen is being tested in phase III clinical trials as treatment for advanced prostate cancer resistant to castration in association with docetaxel. For a detailed analysis of CLU in prostate cancer, see the review by Higano.31

CLU and breast cancer
Table 1 summarizes the findings from various studies of breast cancer discussed below. sCLU overexpression was associated with resistance to preoperative neoadjuvant chemotherapy in primary breast cancer.35

In breast cancer, the situation may be paralleled to that of prostate: the research by Wang et al45 suggests that sCLU confers to breast cancer cells resistance to TNF-α–induced apoptosis through NF-κB activation and Bcl-2 overexpression.

Niu et al46 investigated the effects of sCLU inhibition with small interfering RNAs (siRNAs) on cell motility, invasion, and growth in vitro and in vivo in breast cancer cells. MDA-MB-231 cells were transfected with pSuper-siRNA/sCLU. The results showed that sCLU silencing significantly inhibited the proliferation of MDA-MB-231 cells. The invasion and migration ability were also dramatically decreased. They also demonstrated that sCLU silencing could increase the apoptosis rate of cells, resulting in the inhibition of cell growth. The effects of sCLU silencing on tumor growth and metastatic progression in an orthotopic breast cancer model showed that orthotopic primary tumors derived from MDA-MB-231/pSuper sCLU siRNA cells grew significantly slower than tumors derived from parental MDA-MB-231 or MDA-MB-231/pSuper scramble siRNA cells, and metastasized less to the lungs. These data suggest that sCLU plays a significant role in tumor growth and metastatic progression in certain breast cancer cell lines. The authors consider that knocking down sCLU gene expression may provide a valuable method for breast cancer therapy.

sCLU knockdown in the MDA-231 breast cancer cell line reduced cell migration and invasion in vitro and decreased metastatic potential.37

Triple-negative breast cancer patients were positive for sCLU in a remarkably higher percentage than were patients who were not triple negative, based on immunohistochemical analysis, and sCLU expression. In this triple negative subtype, a larger tumor size, an axillary node status, and a higher clinical stage were observed. Furthermore, they used a spontaneous breast cancer mouse strain with a triple-negative genotype to detect the sCLU dynamic expression in breast cancer oncogenesis. The sCLU mRNA and protein expression in the tumor and hyperplastic epithelium were upregulated and reached a peak compared with those of a normal mammary gland. These results suggest that sCLU is involved in the initiation of triple-negative breast cancer, such that a clinical trial of an anti-CLU treatment for triple-negative breast cancer would be beneficial.46

A previous study47 indicated that CLU expression was associated with estrogen and progesterone receptor-negative status and with the progression from primary carcinoma to metastatic carcinoma in lymph nodes. The study also described correlation between CLU expression and tumor size.

In a Phase II trial concluded in 2009 of custirsen in combination with docetaxel in metastatic breast cancer, custirsen was well tolerated and clinical activity was recorded, but the

Table 1 Summary of findings in breast cancer

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<td>sCLU overexpression is associated with resistance to preoperative neoadjuvant chemotherapy</td>
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<td>45</td>
<td>sCLU confers breast cancer cells resistance to TNF-α apoptosis</td>
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<tr>
<td>36</td>
<td>sCLU silencing significantly inhibited the proliferation of MDA-MB-231 cells, and decreased dramatically invasion and migration</td>
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<td>37</td>
<td>sCLU knockdown in MDA-231 cells reduced cell migration and invasion and decreased metastatic potential</td>
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<td>Overexpression of sCLU is seen in higher percentage in triple negative breast cancer</td>
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<td>47</td>
<td>sCLU expression is associated with negative estrogen and progesterone receptor status</td>
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<td>45</td>
<td>Clusterin inhibition enhanced paclitaxel sensitivity</td>
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<td>48</td>
<td>Custirsen did not show advantages when associated with chemotherapy of breast cancer</td>
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<td>38</td>
<td>Clusterin overexpression enhances metastatic progression</td>
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Abbreviations: sCLU, secretory clusterin; TNF, tumor necrosis factor.
number of responses was insufficient to meet the criteria set for accrual of a second stage. The 15 patients included in this study were all positive for estrogen receptors, progesterone receptors, or both. No triple negatives were recorded. This research had very few patients and did not include triple-negative cases, so according to our view the results may be biased. In another study, clusterin inhibition in vivo and in vitro enhanced paclitaxel chemosensitivity in breast cancer cells.

**CLU and colorectal cancer**

Inhibition of thrombospondin-1 and clusterin abolished p38 mitogen-activated protein kinase phosphorylation, MMP-9 activity, and platelet-stimulated colon cancer invasion. Platelet-secreted thrombospondin-1 and clusterin promote the signal regulation of MMP-9 in platelet-induced colonic cancer invasion via a p38 mitogen-activated protein kinase-regulated pathway. These findings are relevant to the development of therapeutic approaches preventing and reducing tumor cell metastasis induced by colon adenocarcinoma.

In the colon, overexpression of the sCLU was found only in the cytoplasm of highly infiltrating tumors and metastatic nodes. The increased level of the secreted form and the disappearance of the nuclear unglycosylated isoform were directly connected to increased cell survival, aggressiveness, and enhanced metastatic potential.

Normal colonic mucosal cells depend on CLU for survival when a stressor condition appears, such as butyrate-induced cell death.

**CLU and ovarian cancer**

Increased clusterin expression is correlated with more aggressive biologic behavior and impaired survival in ovarian cancer. Promoter methylation of the CLU gene can be readily detected, although at low frequencies, in ovarian epithelial tumors and is significantly associated with decreased protein expression of the gene.

Cyclic adenosine monophosphate (cAMP) induces apoptosis in granulosa cells of rat and human ovary by unidentified mechanisms. The apoptotic effect of cAMP was accompanied by an increase in the expression of p53 and Bax proteins. No change in Bcl-2 protein level was observed in cAMP-treated granulosa cells as compared to controls. These data suggest that cAMP may activate apoptosis in granulosa cells by shifting the ratio of the death promoter genes to death repressor genes via alteration of p53 and Bax expression. cAMP was also shown to inhibit CLU gene expression, suggesting a role for this protein in cAMP-induced apoptosis in granulosa cells.

Treatment of granulosa cells with an antisense oligonucleotide against CLU resulted in an increase in the apoptotic cell death compared to the control. These findings indicate that depletion of CLU can lead to the programmed cell death in the ovary, suggesting a functional role for this protein in follicular atresia.

**CLU and pancreatic cancer**

CLU induction accompanies pancreas regeneration in the rat. CLU induction was accompanied in the regenerating pancreas by a robust development of new lobules with ductules, acini, and endocrine islets after partial pancreatectomy. In CLU knockout mice, however, pancreatectomy resulted in a poor formation of regenerating lobules. In particular, regeneration of beta cells was also significantly reduced and was associated with persistent hyperglycemia. Duct cells obtained from pancreatectomized CLU knockout mice exhibited impaired beta-cell formation in vitro; this was restored by administration of exogenous CLU. These findings suggest that CLU plays a critical role to promote both exocrine and endocrine regeneration following pancreatic injury, as well as for in vitro beta-cell regeneration, so that CLU is an important factor in pancreatic growth.

Transfection of clusterin cDNA into primary cultured pancreatic duct cells resulted in a 2.5-fold increase in cell proliferation and induced transformation of nondifferentiated duct cells into differentiated cells displaying cytokeratin immunoreactivity. These results confirm that CLU may play essential roles in the neogenic regeneration of pancreatic tissue by stimulating proliferation and differentiation of duct cells.

Knockdown of CLU sensitizes pancreatic cancer cells to gemcitabine chemotherapy by ERK1/2 inactivation.

**Table 2 Summary of findings in renal cell carcinoma**

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<td>Introducing the CLU gene into human renal cell carcinoma enhances their metastatic potential</td>
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<td>CLU inhibition using custirsen synergistically enhanced antitumor activity of sorafenib</td>
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<td>34</td>
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<tr>
<td>32</td>
<td>Knockdown of CLU inhibits the growth and migration of renal carcinoma cells</td>
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**Abbreviation:** CLU, clusterin.
CLU and hepatocarcinoma
Hepatocellular carcinoma (HCC) is one of the most common and aggressive tumors. The prognosis of HCC is extremely poor with a five-year survival rate of less than 5% for late primary HCC. Despite recent advances in the diagnosis and treatment of HCC, the mortality rate of HCC remains high and is mainly attributed to the high rate of intrahepatic metastasis after treatment.60

High levels of sCLU expression have been associated with migration, invasion, and metastasis, including HCC.61 It has been shown that CLU is overexpressed in metastatic human HCC tissue compared to nonmetastatic HCC tissue,61 implicating a role for CLU in HCC progression.

Recent studies showed that downregulation of CLU sensitizes cells to chemotherapy and radiotherapy, not only in prostate cancer but also in other malignant tissues like the lung and breast.52,45 In hepatocellular carcinoma, it was not clear whether CLU silencing inhibited the invasion and metastasis, but Chen et al60 provided evidence that CLU plays an important role in HCC invasiveness by increasing MMP-2 expression and decreasing E-cadherin expression.

Lau et al61 showed that overexpression of CLU in hepatoma cells transfected with CLU genes increased cell migration by twofold in vitro and formation of metastatic tumor nodules in the liver by eightfold in vivo. sCLU overexpression contributes to oxaliplatin resistance by activating the AKT pathway in hepatocellular carcinoma.64

CLU in renal cell carcinoma
Introducing the CLU gene into human CCRC enhances the metastatic potential (Table 2).65 We mentioned previously that CLU inhibition using custirsen synergistically enhanced the antitumor activity of sorafenib in a human CCRC model.33

CLU and endometrial and cervical cancer
Estrogen increases the paclitaxel resistance of endometrial cancer cell lines by increasing CLU expression.66 CLU confers paclitaxel resistance in cervical cancer.67

CLU and lung cancer
Yan et al66 suggested that sCLU plays a positive role in non-small-cell lung cancer (NSCLC) cell proliferation, which may be mediated by the PI3K/AKT signaling pathway, and demonstrated that RNA interference-mediated sCLU gene silencing may provide a novel therapeutic strategy in the treatment of NSCLC because it reduced tumor cell proliferation, blocked the PI3K/AKT pathway, and induced apoptosis.

Although CLU expression has been associated with various human malignancies, the mechanisms by which CLU promotes cancer progression and metastasis have not been fully elucidated. Chou et al63 found that CLU regulates epithelial-to-mesenchymal transitions and aggressive behavior of human lung adenocarcinoma cells through modulating ERK signaling and Slug/E-cadherin expression.

In small cell lung cancer cell lines, oblimersen, an antisense Bcl-2 oligodeoxynucleotide, has increased response to radiation in vivo and in vitro. The mechanism of action of oblimersen resembles that of custirsen in the sense of increasing apoptosis.69

CLU and melanoma
Hoeller et al70 reported that in tissue samples, CLU expression was low in nevi but high in primary melanoma and melanoma metastases. CLU was also strongly expressed in melanoma cell lines but was barely detectable in cultured melanocytes. To elucidate a possible role of CLU in melanoma drug resistance, CLU expression was regulated by either plasmid-driven overexpression or by antisense-mediated downregulation. CLU overexpression was associated with an increase in drug resistance, ie, with an increased survival of melanoma cells in the presence of cytotoxic drugs. In contrast, downregulation of CLU by antisense oligonucleotides directed against CLU mRNA significantly reduced drug resistance.

Shannan et al71 studied the expression of CLU immunohistochemically in primary cutaneous malignant melanomas, metastasis of melanomas, and acquired melanocytic nevi. CLU immunoreactivity was found in the two former cases in contrast to acquired nevi. Treatment with vitamin D was observed to modulate CLU expression in vitamin-D–responsive cells but not in resistant melanoma cell lines.

CLU and head and neck squamous cell carcinoma
Overexpression of CLU in head and neck squamous cell carcinoma has been reported, but no relationship implications were made.72

Early versus late inhibition of CLU
There are certain evidences that inhibition of sCLU in late stages of tumor evolution may have beneficial effects in therapy, whereas in the early stages, effects may be the opposite.

In vitro, CLU silencing by antisense oligonucleotides and siRNAs directed against CLU mRNA in CLU-rich lung cancer cell lines sensitized cells to chemotherapy and radiotherapy and decreased their metastatic potential. In vivo, a recent work analyzed the prognostic role of CLU in NSCLC,
showing that CLU-positive patients with lung cancer had a better overall survival and disease-free survival than those with CLU-negative tumors. These data are contradictory to the promising in vitro results.

Based on this, Panico et al. theorize that in early-stage lung cancers CLU represents a positive biomarker correlating with better overall survival. In patients with advanced cancer, already treated with chemotherapy and radiotherapy, the induction of CLU may confer resistance to the treatments.

Summary and future perspectives
The targeting of the apoptotic pathway as a new cancer treatment modality is in accelerated development with novel drugs like oblimersen (a Bcl-2 antisense oligonucleotide blocker), and a combination of apaterson (OGX-427, an antisense oligonucleotide that targets HSP27; OncoGenex Pharmaceuticals Inc.), gemcitabine, and cisplatin is currently being investigated in a Phase II trial of patients with advanced bladder cancer.

CLU has been identified as a valid target for cancer treatment, not only because of its antiapoptotic actions after chemotherapy, but also for other protumor effects it can produce, like PI3K/AKT pathway activation, vascular endothelial growth factor increase, ERK1/2 signaling, MMP-9 overexpression, and NF-kB regulation. CLU inhibition using custirsen synergistically enhances heat-shock protein 90 inhibitor activity by suppressing the heat-shock response in castrate-resistant prostate cancer.

The possible future of custirsen, once it is approved by the US Food and Drug Administration for castration-resistant prostate cancer with metastasis, will be an extension of indications to more tumors. Preclinical evidence warrants the testing of custirsen in CCRC, breast cancer (specifically in triple-negative cases), pancreatic adenocarcinoma, and hepatocarcinoma. In the case of breast cancer, the work by Zhang et al. and Redondo et al. warrants further research of custirsen in spite of the lack of evidence of clinical benefit in the research by Chia et al.

There are certain clues that indicate that inhibition of CLU is a valid therapy in advanced cancer, but this may not be true in early cases. Thus, a major issue in cancer evolution that remains unanswered is if CLU represents a driving force of tumorigenesis or a late phenomenon after chemotherapy. According to the literature here reviewed, we are inclined to think of clusterin as a protein that is overexpressed after intensive chemotherapy and plays a key role in the antiapoptotic defenses against cellular stress induced by chemotherapy. Consequently, CLU inhibition should probably not be attempted in early-stage treatment of cancer.

Disclosure
The author reports no conflicts of interest in this work.

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