Human Antibody Fusion Proteins/Antibody Drug Conjugates in Breast and Ovarian Cancer

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Summary
Considerable research efforts have been dedicated to understanding ovarian and breast cancer mechanisms, but there has been little progress translating the research into effective clinical applications. Hence, personalized/precision medicine has emerged because of its potential to improve the accuracy of tumor targeting and minimize toxicity to normal tissue. Targeted therapy in both breast and ovarian cancer has focused on antibodies, antibody drug conjugates (ADCs), and very recently the introduction of human antibody fusion proteins. Small molecule inhibitors and monoclonal antibodies (mAbs) are used in conjunction with chemotherapeutic drugs as a form of treatment but problems arise from a board expression of the target antigen in healthy tissues. Also, insufficient tumor penetration due to tight binding affinity and macromolecular size of mAbs compromise the efficacy of these ADCs. A more targeted approach is thus needed, and ADCs were designed to meet this need. However, in ADCs the method of conjugation of drug to antibody is >1, altering the structure of the drug which leads to off-target effects. Random conjugation also causes the drug to affect the pharmacokinetics and biodistribution of the antibody and may cause nonspecific binding and internalization. Recombinant therapeutic proteins achieve controlled conjugation reactions and combine cytotoxicity and targeting in one molecule. They can also be engineered to extend half-life, stability and mechanism of action, and offer novel delivery routes. SNAP-tag fusion proteins are an example of a theranostic recombinant protein as they provide a unique antibody format to conjugate a variety of benzyl guanine modified labels, e.g. fluorophores and photosensitizers in a 1:1 stoichiometry. On the one hand, SNAP tag fusions can be used to optically image tumors when conjugated to a fluorophore, and on the other hand the recombinant proteins can induce necrosis/apoptosis in the tumor when conjugated to a photosensitizer upon exposure to a changeable wavelength of light. The dual nature of SNAP-tag fusions as both a diagnostic and therapeutic tool reinforces its significant role in cancer treatment in an era of precision medicine.

Introduction
Breast cancer is the most commonly diagnosed cancer type among women, and ovarian cancer is the sixth most commonly diagnosed cancer in the world [1, 2]. Both types of cancers are strongly associated with mutations in the tumor suppressor genes BRCA1 and BRCA2 [3] and the HER2/neu proto-oncogene [1, 4] and are linked to mutations in genes associated with other inherited autosomal disorders such as Li-Fraumeni (TP53), Peutz-Jeghers (STK11/LKB1) and Cowden syndrome (PTEN) [5]. Breast can-
Antibodies in Breast and Ovarian Cancer

The main difference between ADCs and fusion proteins is attributed to the fact how they are generated. In ADCs, non-cleavable linkers or chemically labile linkers, including acid-cleavable linkers and reducible linkers, help conjugate antibody to drug [29–31]. Due to the chemical method of conjugation, most of these ADCs exist as homogeneous mixtures, which can result in a narrow therapeutic window and have major pharmacokinetic implications [32]. Fusion proteins are made from a fusion gene which is created by joining parts of two different genes together; as a result there is no chemical conjugation necessary, and the constructs are homogeneous [33]. Both these therapeutic agents harness the specific nature of the mAb as an effective means of targeting the diseased cell. In ADCs, the goal is for the antibody to bind to the antigen on the cancer cell, become internalized, and then release the toxic payload to the cancer cell. Linkers are designed to break inside the cancer cells when exposed to a specific pH or a proteasome [34, 35]. In fusion proteins, the goal is to guide the ligand specifically to tumor sites using the mAb linked to a protein moiety [29]. The variable regions (Fv) of antibodies or single-chain variable fragments (scFvs) are generally genetically fused to the protein moiety. scFvs are composed of variable-heavy (VH) and variable-light (VL) domains connected via a glycine, serine, or threonine linker to either the N-terminus of the VH or the C-terminus of the VL [36, 37]. Numerous research groups have constructed recombinant antibody–cytokine fusion proteins composed of the ligands IL-2, IL-12, IL-21, TNF-α, and IFN-α, IFN-β, and IFN-γ which have shown anti-tumor activity [29, 38–42]. Affiblceptor (VEGF-Traps) is a fusion protein which combines ligand-binding elements taken from the extracellular components of the VEGF receptor 1 (VEGFR-1) and VEGFR-2 fused to the Fc portion of IgG [43, 44]. A recent study revealed a greater efficacy for affiblceptor than nesvacumab (an anti-Ang2 antibody) in neoadjuvant/adjuvant chemotherapy for TNBC [45].

The TRAIL-based fusion protein Meso-T3 was found to selectively accumulate on MUC16-expressing cancer targets and increase cytotoxic activity both in vitro and in vivo [46]. Compared to non-targeted TR3, Meso-T3 displayed a much reduced killing potency on cells that lacked MUC16 [47]. The human HIV-1 TAT interactive protein 2 (HTATIP2/TIP30) is an evolutionarily conserved protein...
Antibody-Drug Conjugates and Fusion Proteins in Breast and Ovarian Cancer

At least 2 ADCs for breast cancer are in advanced stages of development: sacituzumab govitecan and glembatumumab vedotin. Sacituzumab govitecan is a conjugate of the humanized anti-Trop-2 monoclonal antibody linked with SN-38, the active metabolite of the chemotherapeutic agent irinotecan [49, 50]. SN-38 is too toxic to administer directly to patients, but linkage to an antibody allows the drug to specifically target cells containing Trop-2 [49, 50]. The fully human IgG2 monoclonal antibody glembatumumab is linked to the mitotane inhibitor called monomethyl auristatin E and targets cancer cells expressing the transmembrane glycoprotein NMB. When patients with metastatic breast cancer were randomly assigned to glembatumumab vedotin, survival rates for metastatic breast cancer patients were significantly better with glembatumumab vedotin treatment than with standard treatment [51]. One ADC that has already received FDA approval is ado-trastuzumab emtansine (T-DM1; Kadcyla) which consists of the monoclonal antibody trastuzumab linked to the cytotoxic agent emtansine [52]. The phase III THERESA study revealed that physicians’ choice of this ADC improved overall survival in patients with previously treated HER2+ metastatic breast cancer [52].

In platinum-resistant epithelial ovarian cancer, the ADC called nirvutaxel soravatserine (IMGN833) is being investigated and consists of a monoclonal antibody that binds to the folate receptor-alpha and is linked to a tubulin-disrupting maytansinoid DM4 chemotherapeutic agent [53]. TIM-1 expression is also upregulated in ovarian carcinomas. An ADC was produced with the anti-TIM-1 antibody covalently linked to monomethyl auristatin E (MMAE) and designated CDX-014 [54]. Other ADCs include PGT-0667020 which consists of a humanized monoclonal antibody against the PTK7 antigen linked to the microtubule inhibitor auristatin. DMOT4059A, a humanized anti-mesothelin mAb conjugated to the antibiotical agent MMAE, was found to have a tolerable safety profile and antitumor activity in both pancreatic and ovarian cancer [55, 56]. In 2016, a potent ADC called XMT-1536 was discovered. It consists of XMT-1535, a novel humanized anti-Trop-2 monoclonal antibody linked with SN-38, the active metabolite of the microtubule inhibitor called monomethyl auristatin E (MMAE) and designated CDX-014 [54]. One ADC that has already received FDA approval is ado-trastuzumab emtansine (T-DM1; Kadcyla) which consists of a monoclonal antibody that binds to the folate receptor-alpha and is linked to a tubulin-disrupting maytansinoid DM4 chemotherapeutic agent [53]. TIM-1 expression is also upregulated in ovarian carcinomas. An ADC was produced with the anti-TIM-1 antibody covalently linked to monomethyl auristatin E (MMAE) and designated CDX-014 [54]. Other ADCs include PGT-0667020 which consists of a humanized monoclonal antibody against the PTK7 antigen linked to the microtubule inhibitor auristatin. DMOT4059A, a humanized anti-mesothelin mAb conjugated to the antibiotical agent MMAE, was found to have a tolerable safety profile and antitumor activity in both pancreatic and ovarian cancer [55, 56]. In 2016, a potent ADC called XMT-1536 was discovered. It consists of XMT-1535, a novel humanized anti-NaPi2b antibody conjugated to 15 MMAE molecules. In an ovarian cancer xenograft model, a single dose (5 mg/kg) of XMT-1536 induced complete tumor regression [57].

Despite significant advances in the field of ADCs, lack of specificity to the target antigen still remains a problem due to the method of drug to antibody conjugation. A drug can bind to multiple side chains on the antibody such that not only does the configuration of each conjugate differ, but so does the number of drug molecules attached to each [58]. As a result, the pharmacokinetic properties and efficacy of each ADC differs [59, 60]. Despite the more controlled conjugation strategy afforded by recombinant fusion proteins, the intrinsic challenges that need to be overcome include immunogenicity that may occur due to the formation of novel epitopes at the junction between the fusion partners even if fully human proteins are connected [61]. SNAP-tag, when genetically fused to scFvs, creates a single conjugation site on the antibody fragment for the site-specific binding of variety of molecules. The SNAP-tag was pioneered by Kai Johnsson, and is derived from the human DNA repair enzyme O(6)-alkylguanine-DNA alkyltransferase [62, 63]. It reacts with para-substituted O6-benzylgua-nine (BG) derivatives by transferring the substituted benzyl group to its active site through a nucleophilic substitution reaction while releasing free guanine [64]. Figure 1 shows a photosensitizer conjugated to SNAP.

This achieves efficient covalent conjugation to any BG-modified substrate under physiological conditions with a 1:1 stoichiometry, generating a homogeneous immunotherapeutic agent [64, 65]. A SNAP-tag fusion protein is composed of the SNAP protein co-expressed with a tumor-specific antibody fragment. The expressed fusion protein is highly selective, site-specific, and resistant due to its high affinity for an array of BG substrates, e.g. toxins, fluorophores or photosensitizers [6–66]. The conjugation of photosensitizers to SNAP-tag has made it an indispensable tool in photodynamic therapy.

**Photoimmunotherapy**

Photodynamic therapy (PDT) has long been the gold standard of treatment for cancer and uses a drug called a photosensitizer followed by local illumination with visible light of specific wavelength(s) to induce apoptosis or necrosis in a tumor [67]. One of the greatest challenges in PDT is the lack of specificity. To overcome this challenge, photoimmunoconjugates have been designed that deliver the photosensitizer directly to the tumor site [68]. These conjugates consist of photosensitizers linked to tumor-spe-
specific mAbs or scFvs in an approach known as photoimmunotherapy (PIT) [68]. This treatment modality is enhanced by imaging in the near infrared (NIR) spectral region. Owing to reduced photon scattering, light absorption and auto-fluorescence as well as increased light penetration into tissue, NIR-PIT is a newly developed cell-selective cancer therapy [69]. Recently, IRDye®700DX N-hydroxysuccinimide ester (IR700) has shown to be a promising photosensitizer because of its high purity and photosensitivity. The dye has a strong absorption peak close to 700 nm allowing light to penetrate deep into the tissue [70, 71]. Moreover, IR700 does not develop off-target effects since they cannot be taken up by cells and cannot exert cytotoxicity even after illumination [72, 73].

To date, a range of research studies have investigated the use of IR700 in PIT. The antibody fragment (scFv-425) that binds to EGFR was used as a model to investigate the use of SNAP-tag fusions as an improved conjugation strategy targeting skin, breast, pancreatic and ovarian cancer cells. The scFv-425-SNAP-fusion protein was conjugated to a range of BG-modified probes: a microtubule inhibitor auristatin F (AURIF) [74], a chlorin 6 photosensitizer in PDT [75], and the IR700 fluorophore in PIT [76, 77]. All these probes were specifically targeted to the tumor and induced cytotoxicity. For example, in the case of cancerous ovarian cells, the SNAP fusion protein was able to eliminate EGFR+ cells with IC₅₀ values of 45–66 nmol/l and 40–90 nmol/l [77]. In the case of breast cancer cells, the recombinant fusion proteins were stable in human serum, were internalized rapidly, and demonstrated potent inhibitory and pro-apoptotic effects in vitro at 3–21 nmol/l [74].

In another study, mAbs targeting EGFR were conjugated to IR700 and irradiated with NIR light, and in vivo tumor shrinkage was observed [73]. The choice of the mAb in photosensitizer conjugates may also influence the effectiveness of PIT. IR700 was conjugated to either cetuximab (cet-IR700) or panitumumab (pan-IR700), and their efficacies were evaluated in EGF spheroids [78]. Although cet-IR700 and pan-IR700 showed identical in vitro characteristics, pan-IR700 showed better therapeutic tumor responses than cet-IR700 in vivo models due to the prolonged retention of the conjugate in circulation. This also suggests that retention in the circulation is advantageous for tumor responses to PIT [78]. Recently, a promising ADC for the treatment of mesothelin-expressing tumors was discovered. It is composed of a humanized mesothelin antibody, known as hYP218, conjugated to IR700 [79]. The hYP218-IR700 showed specific binding to mesothelin-expressing cells, and cell-specific killing was observed in vitro [79]. After irradiation with NIR light, the hYP218-IR700 demonstrated high tumor accumulation-to-background ratio and significa ntly inhibited tumor growth and prolonged survival compared to the control groups [79]. The first step and most common form of treatment for breast and ovarian cancer is surgery, but the heterogeneous nature of the tumors result in poor tissue demarcation and destruction of normal cells. These challenges in surgery have been improved by NIR fluorescence image-guided surgery which has the potential to improve patient outcome and increase the completeness of surgery by decreasing the morbidity associated with damage to normal structures [80].

### Optical Imaging in Guided Surgery

Types of breast cancer surgery include lumpectomy (removal of only the tumor) and mastectomy (removal of all the breast tissue) [81]. In advanced ovarian cancer, cytoreductive surgery is the accepted form of management even though the 5-year survival rate remains at about 40% [82]. The goal of cytoreductive surgery or debulking is to remove all visible tumors. This surgery is limited considering side effects generated by multivisceral resections, i.e. of the colon, spleen and/or the gallbladder as well as of parts of the stomach, liver, and/or pancreas, and some patients die within 30 days of surgery [83, 84]. Thus, both breast and ovarian cancer surgery is improving a good chance to avoid the accurate discrimination of cancerous tissue leading to radical resections of the tumor. In view of this, imaging modalities, e.g. magnetic resonance imaging (MRI), CT scanning and ultrasound techniques, have played a significant role in the field of surgical image guidance, especially during endoscopy [85, 86].

Improvements in preoperative imaging techniques have made a meaningful impact on cancer patient care [80]. Specialized intraoperative imaging systems for open surgery [87–89], laparoscopy [90, 91], and robotic surgery [92, 93] have emerged. Using these systems, NIR fluorescent contrast agents can be visualized with acquisition times in the millisecond range, enabling real-time guidance during surgery [79]. Development of novel NIR fluorescent contrast agents is dependent on the availability of clinically compatible fluorophores [80]. To date only two fluorophores have been reported to be in the process of clinical translation: IRDye 800CW (LI-COR Biosciences, Lincoln, NE, USA) and ZW800-1 (The FLARE Foundation, Wayland, MA, USA) [80, 94]. Both compounds are small molecules, characterized as non-toxic, and can be conjugated to targeting moieties. They are characterized by having renal and hepatic clearance, enabling these agents to be used for imaging of ureters and bile ducts [80, 95].

Additionally, organic fluorophores can be conjugated to SNAP-tag fusion proteins to generate imaging probes for in vitro and in vivo diagnostics [96, 97]. SNAP-tag imaging probes have advantages over autofluorescent protein-antibody probes. These advantages include improved brightness and increased photosensitivity, no chromophore maturation or impact on the labeled proteins function, and shorter half-life with increased retention time of the probes [98–101]. The tumor retention of the conjugate is highly dependent on its binding affinity and internalization, whilst for novel antibody fragments, this binding affinity varies when forming part of fusion proteins [102]. It is thus essential that the above-mentioned parameters are continuously optimized. To this extent, Niesen et al. [103] recently developed a convenient, SNAP-tag-specific SPR spectroscopy assay to measure the binding affinity and functionality of novel scFv-SNAP-tag fusion proteins for use in future immunoantherapeutic production.

The size of the probe also affects both its ability to penetrate tissue and the rate at which it is cleared from circulation [104]. Because deletions in the SNAP-tag gene sequence have reduced its size to 20 kDa compared to that of the wild-type AGT protein or...
endogenous fluorescent proteins, SNAP-tag has a much higher penetrability than its larger counterparts. It is in fact virtually unrestricted in its ability to access any cellular compartment [99]. The intermediate size of the scFv/SNAP construct (~48 kDa) facilitates rapid accumulation and efficacious tumor binding soon after injection together with rapid renal clearance, while still producing a high tumor-to-background ratio with high tumor visualization and low nonspecific background signal [104]. This is ideal for receptor expression monitoring when imaging is done within a short period and further enhances the tumor signal as excess probe is rapidly cleared from the bloodstream, minimizing background interference. [98, 104–106].

In a study by Gong et al. [106], the NIR fluorescent SNAP-tag substrate BG-800 was synthesized by conjugating an IRDye 800CW to the benzyl-guanine amino group (BG-NH2) of the protein tag. Because BG-800 was cell impermeable, the SNAP-ADRβ2 fusion protein was used in such a way that ADRβ2 directed the localization of SNAP-fusion protein to the cell membrane. BG-800 reacted with SNAP-ADRβ2 in both cell lysate and live cell culture [106]. The tumors expressing SNAP-ADRβ2 was then visualized using BG-800 conjugated to the IRDye 800CW. In another study, rapid optical imaging of EGF receptor expression was monitored with an EGFR-specific 425 (scFv) SNAP fusion protein labeled with the NIR dye BG-747 [104]. The 425 (scFv) SNAP fusion protein accumulated rapidly and specifically at the tumor site. Its small size allowed efficient renal clearance and a high tumor-to-background ratio [104].

Nanobodies and affibodies have recently become tumor imaging agents through conjugation to the IRDye 800CW. Affibodies are derived from the staphylococcal protein A and are attractive for imaging purposes due to their small size and low immunogenicity [107, 108]. Similarly, nanobodies are single-domain antibodies derived from the heavy chain of the camelid family [109]. Unlike mAbs, these fragments do not need to undergo partial unfolding as their hydrophobic patches are adequately exposed to facilitate binding to receptors [110]. An anti-EGFR nanobody 7D12 and cetuximab were conjugated to IRDye 800CW to visualize tumors. 7D12 allowed the visualization of tumors as early as 30 min post injection in comparison to cetuximab [111]. In another similar study, the EGFR-specific affibody (Af800), panitumumab (Pan800), and EGF (EGF800) were labeled with the IRDye 800CW. Highest binding affinities were noticed for Pan800 and af800, and the EGFR tumors generated the highest signals for Pan800 and af800 [112]. These studies prove that nanobodies and affibodies can similarly be co-expressed with SNAP-tag and conjugated to organic fluorophores for imaging of tumors in ovarian or breast cancer, and research in this area is ongoing.

Fluorescence optical imaging also has the advantage of multiple channels which can be employed to image two or more targets simultaneously. The clinical antibodies, cetuximab and trastuzumab were labeled with Cy5.5 and Cy7, respectively [113]. When mice were injected with a cocktail of cetuximab-Cy5.5 and trastuzumab-Cy7, A431 and 3T3/HER2+ tumors could be detected distinctly based on the Cy5.5 and Cy7 spectral images [113]. In a subsequent study three antibodies (cetuximab, trastuzumab and daciluzumab) were labeled with three different fluorophores (Cy5, Cy7 and AlexaFluor700). Spectrally resolved fluorescence imaging showed that these probes clearly distinguished their respective targeting tumors (A431, 3T3/HER2+ and SP2-Tac) based on their distinct optical spectra [114]. These studies complement recent research into dual-color single molecule imaging of SNAP-tag fusion proteins using an optimal dye pair [101]. The labeling was performed on SNAP-EGFR with BG-Dy549 (green) and BG-CF633 (red) [101]. This study demonstrated how a single SNAP-tag fusion protein can be labeled with a selection of differently colored fluorophores without the need to separately clone each and opens the way for a potentially powerful method of visualizing different antigens on one tumor without worrying about tumor heterogeneity.

**Conclusion**

Efforts in the treatment of breast and ovarian cancer will continue to focus on personalizing treatment to the patient and the tumor. Immunotherapy achieves this goal as it blocks the growth of cancer cells by interfering with specific targeted molecules needed for carcinogenesis and tumor growth, and ADCs and fusion proteins are types of immunotherapeutic agents. A plethora of ADCs currently exist to treat ovarian and breast cancer with a few being approved and others still in clinical trials. The only ADC that has currently been approved for metastatic breast cancer is Kadcyla. Human antibody fusion proteins are now emerging therapeutic tools due to their homogeneity in combining functionality along with its applications in imaging and PIT only serve to exemplify the attractive diagnostic and therapeutic potential of fusion proteins in targeted cancer treatment.

**Disclosure Statement**

There are no conflicts of interest.
References


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