Pharmacologic Suppression of B7-H4 Glycosylation Restores Antitumor Immunity in Immune-Cold Breast Cancers

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Despite widespread utilization of immunotherapy, treating immune-cold tumors has proved to be a challenge. Here, we report that expression of the immune checkpoint molecule B7-H4 is prevalent among immune-cold triple-negative breast cancers (TNBC), where its expression inversely correlates with that of PD-L1. Glycosylation of B7-H4 interferes with its interaction/ubiquitination by AMFR, resulting in B7-H4 stabilization. B7-H4 expression inhibits doxorubicin-induced cell death through the suppression of eIF2α phosphorylation required for calreticulin exposure vis-à-vis the cancer cells. NGI-1, which inhibits B7-H4 glycosylation causing its ubiquitination and subsequent degradation, improves the immunogenic properties of cancer cells treated with doxorubicin, enhancing their phagocytosis by dendritic cells and their capacity to elicit CD8+ IFNγ-producing T-cell responses. In preclinical models of TNBC, a triple combination of NGI-1, camisurubicin (a noncardiotoxic doxorubicin analogue) and PD-L1 blockade was effective in reducing tumor growth. Collectively, our findings uncover a strategy for targeting the immunosuppressive molecule B7-H4.

SIGNIFICANCE: This work unravels the regulation of B7-H4 stability by ubiquitination and glycosylation, which affects tumor immunogenicity, particularly regarding immune-cold breast cancers. The inhibition of B7-H4 glycosylation can be favorably combined with immunogenic chemotherapy and PD-L1 blockade to achieve superior immuno-infiltration of cold tumors, as well as improved tumor growth control.

See related commentary by Pearce and Läubli, p. 1789.

INTRODUCTION

Despite the widespread use of immunotherapy, the poor clinical response by immune-cold tumors is a current challenge. Triple-negative breast cancer (TNBC) is the most aggressive mammary carcinoma subtype. Although a PD-L1 inhibitor has been FDA-approved for the treatment of metastatic TNBC, the majority of patients with TNBC show limited responses, particularly when their tumors are “cold” or “noninflamed” (1). In contrast with “hot” tumors, these immune-cold tumors are characterized by a scarcity of T lymphocyte infiltration, commensurate with their failure to elicit anticancer immunity (2). Our endeavor to identify suitable targets for improving anticancer immune responses against TNBC therapy has drawn our attention to B7-H4.

B7-H4, also known as V-set domain containing T-cell activation inhibitor 1 (VTCN1/B7x/B7 homolog4/B7S1) is a type 1 transmembrane protein and belongs to the coinhibitory B7 family ligands (3-5). It regulates T cells and neutrophils as well as macrophages, and its local expression in the tumor has been linked to poor prognosis for ovarian, uterus, gastric, lung, kidney, and breast cancers (6). Although B7-H4 levels are known to be regulated at the post-transcriptional level, a detailed exploration of its post-translational modifications (PTM) and regulation has been elusive, further spurring our interest in this gene product.

Cancer cells are known to evade the immune system through “don’t eat me” signals including β2 microglobulin, CD24, CD47, and perhaps even PD-L1 (7-12). These signals allow malignant cells to escape recognition and engulfment by dendritic cells (DC), thus resulting in a failure to generate antitumor immune responses. Some chemotherapeutic drugs such as anthracyclines and microtubule-destabilizing agents possess an immune-stimulatory function by triggering immunogenic cell death (ICD; ref. 13). ICD is usually characterized by the cell membrane exposure of damage-associated molecular patterns (DAMP) such as calreticulin (CALR) and HSP70 and HSP90, which serve as “eat me” signals to facilitate the recognition of stressed cancer cells by DCs (14). Extensive biochemical analyses have revealed that the translocation of CALR from the endoplasmic reticulum (ER) to the cell surface relies on a particular ER stress signal, the phosphorylation of eukaryotic initiation factor 2α (eIF2α), which is the pathognomonic hallmark of ICD (7, 15). Nevertheless, how CALR exposure may be suppressed to favor immune evasion remains elusive.

In this study, we unravel a regulatory mechanism by which B7-H4 protein turnover is governed by asparagine...
(N)-linked glycosylation. N-linked glycosylation is a co-and post-translational modification important for protein stability, folding, trafficking, and physiologic function (16, 17). Glycosylation of B7-H4 catalyzed by specific glycosyltransferases (STT3A and UGGG1) antagonizes the B7-H4 ubiquitination by E3 ligase autocrine motility factor receptor (AMFR), thereby preventing the degradation of B7-H4 and hence stabilizing the protein. Abundant B7-H4 can inhibit the phosphorylation of eIF2α, CALR exposure, and cancer cell immunogenicity. We designed a strategy for inhibiting the oligosaccharyltransferase (OST) complex that stabilizes B7-H4, thus reducing its expression and enhancing the immunogenicity of TNBC cells in the context of ICD-eliciting chemotherapy and PD-L1-targeting immunotherapy.

RESULTS
Accumulation of B7-H4 in Immune-Cold Breast Cancers Inversely Correlates with PD-L1 Expression

To identify potential therapeutic targets for immune-cold breast cancer, we performed a bioinformatic analysis of proteomic data concerning 100 immune-relevant proteins from 81 patients with breast cancer in The Cancer Genome Atlas (TCGA; ref. 18). Elevated expression of three proteins, including PROM1, B7-H4 (VTCLN1), and MMP7, was detected in the basal (TNBC) subtype of breast cancer compared with other subtypes (Fig. 1A).

Recent studies have subdivided TNBC into four subsets based on the tumor immune microenvironment (TIME): “fully inflamed” (FI), “margin-restricted” (MR), “stroma-restricted” (SR), and “immune desert” (ID). ID and MR are both poorly infiltrating tumors with signatures of fibrosis. The SR subtype is defined by the infiltration of CD8+ T cells in the stroma and low IFN signature. Only the FI TNBC subtype has a proinflammatory microenvironment defined by a type I IFN gene signature, CD8+ T-cell infiltration in the tumor epithelium, and good outcomes (19). We compared gene expression in FI versus non-fully infiltrated tumors (MR+SR+ID). In 38 TNBC patient specimens from the Gene Expression Omnibus (GEO) database (accession number: GSE88847), the mRNAs coding for the aforementioned immune-relevant proteins’ upregulation of B7-H4 expression was associated with the non-fully infiltrated phenotype (Fig. 1B), contrasting with high expression of CD274 (best known as PD-L1) and IDO1 in hot tumors, consistent with another report (19).

We then systematically measured the protein expression levels of PD-L1, PD-L2, B7-H3, and B7-H4 in a panel of 45 breast and 4 ovarian cell lines (Fig. 1C). B7-H3 and PD-L1 but neither B7-H4 nor PD-L2 were expressed by normal breast cells (MCF-10A, MCF010F, 184-B5, and MCF-12A). In contrast, B7-H4 was detected in 60.9% (25/41) of the tested breast cancer cell lines (among which 10 are TNBC cell lines), compared with 10% breast cancer cells for PD-L2 expression and 25% for PD-L1 expression. Intriguingly, the expression of B7-H4 was inversely correlated with PD-L1 expression in these cell lines (Spearman correlation analysis; Fig. 1D; Supplementary Table S1). An IHC array performed on a tissue array of 110 breast invasive ductal carcinomas confirmed the inverse correlation between B7-H4 and PD-L1 expression (Fig. 1E and F). As compared with normal breast tissue, B7-H4 was increased in hormone receptor-positive (ER+/PR+) and triple-negative breast cancers (Fig. 1G). Many groups have reported a negative correlation between B7-H4 expression and T-cell infiltration, including in breast-invasive ductal carcinomas, uterine endometrioid adenocarcinomas, human esophageal squamous cell carcinoma, and clear cell ovarian cancer (20–23), although some reports are controversial (24–26). In our study, we observed a negative correlation between B7-H4 expression and CD8 T-cell infiltration based on the tissue array (Fig. 1H and I). Taken together, our results suggest that B7-H4 is overexpressed in TNBC, especially in immune-cold tumors.

B7-H4 Glycosylation Antagonizes Its Ubiquitination and Stabilizes the Protein

Previous studies reported that B7-H4 mRNA is transcribed in multiple peripheral tissues in both mouse and human, while its protein expression is low in normal tissues (3, 27, 28). This discrepancy between the mRNA and protein levels of B7-H4 in normal tissues suggests a role for post-transcriptional regulation of B7-H4. Intrigued by the accumulation of B7-H4 in TNBC (Fig. 1A, C, and G), we decided to investigate the PTMs of this protein. The electrophoretic mobility of B7-H4 corresponds to a protein of approximately 50 kDa, with additional bands around 40 kDa and 28 kDa (the predicted size of unglycosylated B7-H4), as determined by immunoblot. Protein extracts from HCC1954, MDA-MB-231, and SKBR3 cells were treated with PNGase F, a peptide:N-glycosidase F, revealing that removal of asparagine-linked carbohydrates

Figure 1. Accumulation of B7-H4 is associated with immune-cold breast cancer and reduced PD-L1 expression. A, Proteomic analysis of 59 immune-relevant proteins in TCGA samples of PAM50-defined intrinsic subtypes including 25 basal-like, 29 luminal A, 33 luminal B, and 18 HER2-enriched tumors, along with three normal breast tissue samples. The genes (rows) are sorted according to the difference between the average proteomic level in basal subtype and the average proteomic in the other cancer types. B, Heat maps depicting expression of immune-relevant genes (mRNA) in the bulk tumor in FI (fully infiltrated), SR (stroma restricted), MR (margin restricted) and ID (immune desert) TNBC (n = 37). C, The protein expression of PD-L1, PD-L2, B7-H3, and B7-H4 in 45 breast and 4 ovarian cancer cell lines were measured by immunoblot. D, Expression of PD-L1 and B7-H4 in the indicated cell lines was quantified using ImageLab. Spearman correlation indicates B7-H4 expression is negatively correlated with PD-L1 expression in the test cancer lines (r = 0.6128, P = 1.43 × 10−4). E-I, Tissue array of 110 breast invasive ductal carcinomas (including 46 cases of ER/PR positive, 37 cases of HER2 positive, and 17 cases of TNBC, and 10 adjacent normal tissue specimens) were subjected to IHC. E, Representative paired IHC staining of B7-H4 and PD-L1. F, Statistical analysis of IHC staining indicates B7-H4 expression is negatively correlated with PD-L1 expression in breast cancer tissues (r = −0.480, P = 1.09 × 10−3). G, The B7-H4 staining in the tissue array was quantified on the basis of the subtypes including ER+/PR+, HER2−, TNBC, and normal breast tissue samples. H, Representative paired IHC of B7-H4 and CD8. I, Statistical analysis of IHC staining indicates that CD8 T-cell number is negatively correlated with B7-H4 expression in breast cancer tissues (r = −0.408, P = 1.00 × 10−1).
Regulation of B7-H4 by PTM Orchestrates Tumor Immunogenicity

A

Protein

Normal Luminal A Luminal B HER2 Basal

B7-H4

Low exp

High exp

Color key

-4 0 4 Row Z-score

B

mRNA

Fl MR SR ID

B7-H4

Low exp

High exp

Color key

-4 0 4 Row Z-score

C

Protein

PD-L1 PD-L2 B7-H3 B7-H4 Actin

D

Spearman correlation

P-D-L1

P-D-L2

P-B7-H3

P-B7-H4

Actin

E

Case 1

Case 2

F

Spearman correlation

r = -0.480

P = 1.09 x 10^{-7}

G

B7-H4 staining

Normal HER2 ER PR TNBC

P = 0.012

P = 0.048

P = 0.117

H

Case 1

Case 2

I

Spearman correlation

r = -0.408

P = 1.0 x 10^{-5}
leads to the disappearance of the 40 and 50 kDa bands and an accentuation of the 28-kDa band (Fig. 2A). Thus, differential N-glycosylation leads to the generation of at least three forms of B7-H4 with high glycosylation (approximately 50 kDa), less glycosylation (approximately 40 kDa) and no glycosylation (28 kDa). Next, we examined the protein stability of B7-H4 by means of a pulse-chase assay with cycloheximide, an inhibitor of new protein biosynthesis. The less-glycosylated and unglycosylated forms of B7-H4 had a faster turnover rate (half-life <2 hours) than the highly glycosylated form (half-life >6 hours; Fig. 2B), suggesting that glycosylation enhances the stability of B7-H4 protein.

The proteasome inhibitor MG132 significantly increased the abundance of the nonglycosylated form of B7-H4. This, along with the fact that MG132 increased ubiquitin conjugates of B7-H4 (Fig. 2C), suggested that B7-H4 turnover is regulated by the ubiquitin–proteasome pathway. Next, we examined the impact of glycosylation on B7-H4 stability by treating cells with the N-glycosylation inhibitor tunicamycin, followed by a pulse-chase assay with cycloheximide. As tunicamycin decreased the glycosylation of B7-H4, it increased the relative abundance of nonglycosylated B7-H4 (Fig. 2D and E), while simultaneously enhancing the turnover of B7-H4. The effect of glycosylation on B7-H4 ubiquitination was confirmed in two additional breast cancer cell lines by immunoblot (Supplementary Fig. S1A) and the ubiquitination assay (Fig. 2F; Supplementary Fig. S1B), showing that the inhibition of N-glycosylation enhanced B7-H4 ubiquitination.

Identification of Glycosylation Sites on B7-H4

To identify the glycosylation and ubiquitination sites on hB7-H4, FLAG-tagged hB7-H4 protein was ectopically expressed in 293T cells, then purified by the affinity capture approach in the absence or presence of PNGase F, followed by mass spectrometry. This led to the identification of a panel of N-linked glycosylation sites at asparagine (N) residues N112, N140, N156, N160, and N255 of B7-H4 (NP_078902.2), as well as that of two ubiquitination sites at lysine (K) residues K146 and K138 (Fig. 2G; Supplementary Fig. S1C and S1D). These two ubiquitination sites were only detected in the presence of PNGase F, indicating that N-linked glycosylation at the asparagine sites may interfere with ubiquitination at the lysine residues. In the next step, we mutated the aforementioned glycosylation sites. Mutation of the five identified asparagines to glutamine residues in B7-H4 (SNQ, amino acids 112, 140, 156, 160, and 255) partially reduced B7-H4 protein glycosylation, as reflected by its shift to higher electrophoretic mobility (Fig. 2H). We also mutated all 16 asparagine residues found in B7-H4 to glutamine residues (16NQ: amino acids 47, 112, 119, 140, 142, 156, 160, 190, 196, 202, 205, 216, 220, 221, 229, and 255), confirming that this manipulation completely blocked B7-H4 glycosylation (Fig. 2H).

Next, cycloheximide chase experiments were performed with these mutants, revealing that hB7-H4–16NQ had an increased turnover rate compared with that of B7-H4 wild-type or other mutants (Supplementary Fig. S1E). The 16NQ mutation also led to a marked increase in ubiquitinated B7-H4 compared with B7-H4 wild-type, in both the absence or presence of MG132 treatment (Supplementary Fig. S1F).

B7-H4 Protein Stability

In the next step, we sought to determine the ubiquitin E3 ligases and glycosyltransferases affecting B7-H4 stability. A FLAG-B7-H4 construct was stably expressed in MDA-MB-468 cells and the B7-H4 protein was purified by affinity capture purification (Fig. 3A), followed by mass spectrometrometric identification of the B7-H4 interactome. Within this interactome, we found the ubiquitin E3 ligase autocrine motility factor receptor (AMFR) and several glycosyltransferases such as UDP-glucose:glycoprotein glucosyltransferase (UGGG1), several OST complex subunits such as...
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**A**

PNGase F

<table>
<thead>
<tr>
<th>B7-H4</th>
<th>HCC1954</th>
<th>MDA-MB-468</th>
<th>SKBR3</th>
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<tbody>
<tr>
<td>High glycosylation</td>
<td>0 h</td>
<td>2 h</td>
<td>4 h</td>
</tr>
<tr>
<td>Less glycosylation</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Nonglycosylation</td>
<td>-</td>
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</table>

Actin

50 kDa

28 kDa

250 kDa

**B**

CHX

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<th>HCC1954</th>
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<th>SKBR3</th>
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</thead>
<tbody>
<tr>
<td>0 h</td>
<td>2 h</td>
<td>4 h</td>
</tr>
<tr>
<td>B7-H4</td>
<td>50 kDa</td>
<td>40 kDa</td>
</tr>
<tr>
<td>Actin</td>
<td>42 kDa</td>
<td>42 kDa</td>
</tr>
</tbody>
</table>

**C**

IP: FLAG

Input

| Ubiquitin-HA | 250 kDa |
| FLAG-hB7-H4 | 250 kDa |

Actin

50 kDa

28 kDa

250 kDa

**D**

DMSO

Tunicamycin

<table>
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<th>4</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>B7-H4</td>
<td>50 kDa</td>
<td>40 kDa</td>
<td>28 kDa</td>
<td>28 kDa</td>
</tr>
<tr>
<td>Actin</td>
<td>42 kDa</td>
<td>42 kDa</td>
<td>42 kDa</td>
<td>42 kDa</td>
</tr>
</tbody>
</table>

**E**

Relative intensity (%)

0 %

150%

**F**

IP: FLAG

Input

| Ubiquitin-HA | 250 kDa |
| FLAG-hB7-H4 | 250 kDa |

Actin

50 kDa

28 kDa

250 kDa

**G**

Overexpression of FLAG-hB7-H4 in 293T cells

Purification of FLAG-hB7-H4

Samples were treated with PNGase F

Mass spectrometry

N112, N140, N156, N160, N255
K146, K138

**H**

| FLAG-hB7-H4 | 50 kDa | 40 kDa | 28 kDa |
| FLAG-K138R | 250 kDa | 50 kDa | 40 kDa |

Actin

42 kDa

**I**

IP: FLAG

| FLAG-B7-H4 | + | - | - | + | - | - | - | - | - |
| FLAG-K138R | + | + | + | + | + | + | + | + | + |
| FLAG-K146R | + | + | + | + | + | + | + | + | + |
| FLAG-K138/146R | + | + | + | + | + | + | + | + | + |
| MG132     | + | + | + | + | + | + | + | + | + |

Ubiquitin

50 kDa

40 kDa

28 kDa

250 kDa

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as STT3A, RPN1, and RPN2 (Fig. 3A), as well as p97, which interacts with AMFR to participate in ER-associated degredation (ERAD). Using a similar experimental approach, the glycosyltransferases RPN1, RPN2, and UGGG1 were confirmed to bind B7-H4 in 293T cells (Supplementary Fig. S2A). Coimmunoprecipitation (co-IP) assays performed on breast cancer cells confirmed the interactions of B7-H4 with AMFR, STT3A, UGGG1, and HSP90 (Fig. 3B). Furthermore, immunostaining followed by confocal microscopy revealed the colocalization of B7-H4 and AMFR or that of B7-H4 and STT3A in the cytoplasm (Fig. 3C), a colocalization that was further corroborated by means of the Duolink proximity ligation assay (Fig. 3D). Of note, the B7-H4 and AMFR expression levels as determined by IHC showed an inverse correlation for an array of 110 specimens of breast invasive ductal carcinomas (r = 0.3464, P = 2.1 × 10−4; Supplementary Fig. S2B and S2C), suggesting a potential role for AMFR in controlling B7-H4 expression. Conversely, B7-H4 and STT3A protein levels were positively correlated with each other in this tissue array (r = 0.3030, P = 1.29 × 10−3; Supplementary Fig. S2B–S2D), in line with a possible role for STT3A in stabilizing B7-H4.

Next, we generated two AMFR knockdown cell lines using two nonoverlapping shRNAs targeting AMFR. AMFR knockdown led to an increase of B7-H4 protein as compared with the vector control (pLKO; Fig. 3E). Transfection-enforced overexpression of AMFR wild-type (AMFR-WT-FLAG) increased B7-H4 ubiquitination, whereas the AMFR mutant with ring-domain mutation C356G/H361A (AMFR-RM-FLAG, which loses its ubiquitin ligase activity; ref. 29) failed to do so, confirming that AMFR functions as a critical E3 ubiquitin ligase for B7-H4 (Supplementary Fig. S2E).

Glycosylation involves several enzymatic steps including several steps involving the OST complex, glucosidases, ER mannosidases, and UGGG1 in the ER (17). The knockdown of the OST catalytic subunit STT3A by two distinct nonoverlapping shRNAs resulted in the reduction of B7-H4 in 293T cells (Supplementary Fig. S2A). Coimmunoprecipitation (co-IP) assays performed on breast cancer cells confirmed the interactions of B7-H4 with AMFR, STT3A, UGGG1, and HSP90 (Fig. 3B). Furthermore, immunostaining followed by confocal microscopy revealed the colocalization of B7-H4 and AMFR or that of B7-H4 and STT3A in the cytoplasm (Fig. 3C), a colocalization that was further corroborated by means of the Duolink proximity ligation assay (Fig. 3D). Of note, the B7-H4 and AMFR expression levels as determined by IHC showed an inverse correlation for an array of 110 specimens of breast invasive ductal carcinomas (r = 0.3464, P = 2.1 × 10−4; Supplementary Fig. S2B and S2C), suggesting a potential role for AMFR in controlling B7-H4 expression. Conversely, B7-H4 and STT3A protein levels were positively correlated with each other in this tissue array (r = 0.3030, P = 1.29 × 10−3; Supplementary Fig. S2B–S2D), in line with a possible role for STT3A in stabilizing B7-H4.

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NGI-1, an aminobenzamide-sulfonamide inhibitor, was initially identified from a cell-based high-throughput screen and lead compound optimization campaign targeting the catalytic subunit STT3A/B of OST. NGI-1 inhibited the glycosylation of B7-H4 and reduced its expression in two breast cancer cells as determined by immunoblot (Fig. 3I). NGI-1 also increased the ubiquitination of B7-H4 (Fig. 3J). NGI-1 was reported to block EGFR signaling in non–small cell lung cancer (NSCLC) and enhance glioma radiosensitivity (31–33). We examined the level of EGFR expression in MDA-MB-468, SKBR3, and mouse breast cancer cells 4T1 cells and E0771 cells by flow cytometry (Supplementary Fig. 3A and B) and immunoblot (Supplementary Fig. 3C), comparing the effect of NGI-1 on B7-H4 and EGFR. We observed that NGI-1 has a modest effect on inhibiting EGFR in HCC1954, MDA-MB-231, and SKBR3 cells, with no effect on EGFR in MDA-MB-468 cells. On the other hand, NGI-1 significantly inhibited B7-H4 in all tested breast cancer cell lines (Supplementary Fig. 3D). EGFR expression is undetectable in 4T1 cells, indicating that 4T1 is an ideal breast cancer cell line to study the effect of NGI-1 on B7-H4 to exclude a possible bias from EGFR. We also compared the effect of glycosylation on the stability of B7-H4 or EGFR by cycloheximide pulse-chase. STT3A knockdown (Supplementary Fig. S3E–S3G) and pre-treatment with 1 μmol/L NGI-1 (Supplementary Fig. S3H–S3J) increased protein turnover of B7-H4 but not EGFR. These experiments indicated that glycosylation affects the turnover of the B7-H4 but not EGFR. In conclusion, the genetic or pharmacologic inhibition of glycosyltransferase STT3A distinguishably increases the E3 ligase AMFR-mediated ubiquitination and degradation of B7-H4.
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Regulation of B7-H4 by PTM Orchestrates Tumor Immunogenicity

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Molecular Analysis of the Interaction between B7-H4 and the E3 Ligase AMFR

To identify the domains of AMFR and B7-H4 responsible for their interaction, we constructed a series of FLAG-tagged B7-H4 and MYC-tagged AMFR truncation mutants (Fig. 4A–D), cotransfected them into cells, and performed co-IP experiments. This approach led to the prediction that the domain comprising amino acid residues 153–241 (corresponding to the Ig-like C2-type domain) in B7-H4 (Fig. 4B) and the RING domain comprising residues 302–379 on AMFR are implicated in the AMFR-B7-H4 interaction (Fig. 4D). Next, we conducted a series of docking and molecular dynamics (MD) simulations. The X-ray structure of the B7-H4 Ig-like V-type domain (PDB: 4GOS; ref. 34) and PD-L1 (PDB: 5JDR; ref. 35), which shares 23.9% sequence identity with B7-H4, were used as templates in homology modeling. The AMFR RING domain (the structure of which has been determined) was docked onto the predicted B7-H4 structure, yielding a molecular model of the interactions between the AMFR RING domain mainly and the B7-H4 Ig-like C2-type domain (Fig. 4E–J). To assess the stability of these complex conformations, we performed MD simulations for the top five clusters among the top 100 ZDOCK docking poses, where the root-mean-square deviations (RMSD) in the atomic positions of AMFR RING domain were evaluated by structurally aligning the generated conformers after superposing their Ig-like C2-type domain. A low RMSD profile over 120-ns simulation time confirmed that the RING domain was highly stable in the bound conformation of the top 1 cluster (Fig. 4F; see the trajectory in Supplementary Movie). Figure 4G and J displays the bound conformation of the top 1 cluster of the ZDOCK docking poses between B7-H4 and AMFR RING domain are shown in (Fig. 4L and N). This suggests that the mutant 16NQ is flexible compared with B7-H4 wild-type in the Ig-like C2-type domain, while the entire structure is still stable.

**Figure 4.** Mapping of molecular domains/motifs and structure-based modeling and simulations reveal critical regions and interfacial interactions involved in the complex formation between B7-H4 and E3 ligase AMFR. A, Schematic diagram of human B7-H4 domains and strategy to engineer a series of B7-H4 deletion mutants. B, Mapping of B7-H4 regions [sequence ranges] involved in interactions with AMFR. The interactions between MYC-AMFR and the displayed FLAG-hB7-H4 fragments were examined by co-IP experiments in 293T cells. C, Schematic diagram of human AMFR domains/motifs and strategy to engineer a series of AMFR deletion mutants. TM, transmembrane domain; RING motif, E3 ligase activity; OS, Oligomerization domain; CUE domain, Coupled Ubiquitin molecules to ER degradation; G2BR, Ube2G2 binding region; VIM, p97/VCP interacting motif. D, Mapping of AMFR domains/motifs that interact with B7-H4. The interactions between MYC-hB7-H4 and the displayed FLAG-AMFR fragments were examined by co-IP experiments in 293T cells. The representative structures of the top 1 cluster of the ZDOCK docking poses between B7-H4 and AMFR RING domain are shown in E. AMFR RING, B7-H4 Ig-like V-type and Ig-like C2-type domains are in cyan, pale green, and green. Ubiquitination sites (K138 and K146) are shown as blue sticks with alpha-carbon atoms highlighted in blue spheres. All asparagines are shown as orange sticks, and the alpha-carbon atoms of five identified asparagines N12, N140, N156, N160, and N255 (not shown in the structure) are highlighted in orange spheres. The two Zn⁺ ions forming coordination bonds are shown as salmon spheres. Residues in the binding interfaces from AMFR and B7-H4 are shown as cyan and green sticks, respectively. Time evolution of the RMSDs of AMFR RING domain in the 120-ns MD simulations of the complex formed with B7-H4 using the start points in E is shown in F. The RMSD was evaluated after structurally aligning the conformers observed during MD trajectories with respect to the B7-H4 Ig-like C2-type domain. The corresponding time evolution of residue-residue interactions between B7-H4 and AMFR RING domain residues are shown in G. Regions shaded in gray refer to time intervals during which the indicated atom pairs (ordinate) made interfacial contacts. The representative structures of the top 2 cluster of the ZDOCK docking poses between B7-H4 and AMFR RING domain are shown in H. Time evolution of the RMSDs of AMFR RING domain in the 120 ns MD simulations of the complex formed with B7-H4 using the start points in H is shown in I. The corresponding time evolution of residue-residue interactions between B7-H4 and AMFR RING domain residues are shown in J. The RMSDs profiles of the B7-H4 Ig-like V-type and C2-type domains are shown in blue and red curves for B7-H4 wild-type and 16NQ mutant, respectively. The corresponding RMSFs values of residues in the two domains are shown in M and N, respectively. The spheres on the red curves indicate the positions of the 16NQ mutations. The RMSDs and RMSFs values for the 16NQ mutant are slightly higher than those of the wild-type in the C2-type domain, although the difference is not statistically significant. Flexibilities (RMSDs and RMSFs) of the Ig-like V-type domain of B7-H4 wild-type and 16NQ mutant are quite similar.

**B7-H4 Inhibits Doxorubicin-Induced ICD through Regulating PERK/eIF2α/CALR Axis**

Immunogenic cell death is defined by the release and exposure of DAMPs from and on dying tumor cells, thus initiating an anticancer immune response (13). One major DAMP is CALR, a protein that usually locates to the lumen of the ER but then translocates to the cell surface when ICD is induced and acts as an “eat-me” signal to facilitate the phagocytosis of portions of the tumor cell by immature DCs (14, 36). The phosphorylation of eukaryotic translation

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**References:**

1. Song et al. (2020). Cancer Discovery, 18(12), 1840-1854. doi: 10.1158/2159-8290.CD-20-0402

2. Cancer Research. (2020). Published OnlineFirst September 16, 2020; DOI: 10.1158/2159-8290.CD-20-0402

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Regulation of B7-H4 by PTM Orchestrates Tumor Immunogenicity

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initiation factor 2 subunit alpha (eIF2α) is a critical event in the molecular cascade leading to CALR exposure on the plasma membrane (15). The B7-H4 interactome includes ER chaperones as well as eIF2α (Fig. 3A; Supplementary Fig. S2A), suggesting a possible connection between B7-H4 and ICD. Results from clinical studies unveiled a critical role for ICD with regard to the success of chemotherapy with anthracyclines such as doxorubicin (13, 37). Camisrubicin (GPX-150) is a noncardiotoxic doxorubicin analogue (38). We observed that doxorubicin-induced or camisrubicin-induced cell death, reduced cell viability, and clone formation of MDA-MB-468 breast cancer cells was not affected by NGI-1 (Supplementary Fig. S5C, S5H, and S5I). Of note, NGI-1 significantly enhanced the doxorubicin-induced phosphorylation of eIF2α, suggesting a possible connection between B7-H4 and ICD.

To investigate how B7-H4 affects eIF2α phosphorylation, we treated FLAG-hB7-H4–expressing cells with doxorubicin and/or NGI-1 and then immunoprecipitated the flagged eIF2α kinase 3 (eIF2AK3, best known as the ER stress kinase PERK), CALR, HSP70, and HSP90. This experiment revealed that B7-H4 interacts with eIF2α, PERK, CALR, and HSP90. Intriguingly, eIF2α was released from the B7-H4 protein complex following doxorubicin treatment (Fig. 5F).

To further investigate the alteration of B7-H4 protein complex in the presence or absence of doxorubicin, we conducted an Iodixanol (OptiPrep) gradient analysis. Doxorubicin induced the formation of novel molecular complex peak (peak II) including eIF2α, eIF2α kinase 3 (eIF2AK3), and eIF2α kinase 1 (eIF2AK1), best known as the ER stress kinase PERK, CALR, HSP70, and HSP90. This experiment revealed that B7-H4 interacts with eIF2α, PERK, CALR, and HSP90. Intriguingly, eIF2α was released from the B7-H4 protein complex following doxorubicin treatment (Fig. 5F).

Taken together, these results suggest that B7-H4 can bind to eIF2α, thus inhibiting its phosphorylation by PERK. Upon the treatment with doxorubicin, eIF2α is released from this inhibitory interaction, allowing for its phosphorylation and the subsequent translocation of DAMPs to the plasma membrane.

**B7-H4 Subverts Spontaneous and Doxorubicin-Induced Immunosurveillance**

In the context of ICD, surface-exposed CALR, HSP70, and HSP90 are potent “eat-me” signals for DCs, thus facilitating the presentation of tumor-associated antigens to CTLs. Moreover, the coculture of malignant cells succumbing to ICD and immature DCs induces DC maturation. We wondered...
Correlated with p-eIF2α staining of B7-H4 and phospho-eIF2α (Ser51) in tissue array BC081120. Statistical analysis of IHC staining indicates B7-H4 expression is negatively correlated with p-eIF2α expression in breast cancer (r = −0.249, P = 8.71 × 10⁻³). D, MDA-MB-468-FLAG-hB7-H4 were treated in the presence or absence of doxorubicin (10 μmol/L) and/or NGI-1 (10 μmol/L) for 24 hours. Then FLAG-hB7-H4 was immunoprecipitated followed by immunoblot. The indicated proteins were examined.

**Figure 5.** (Continued) C and D, SKBR3, MDA-MB-468, MDA-MB-468-vector, and MDA-MB-468-B7-H4 knockout cells were treated with 1 or 10 μmol/L doxorubicin and/or 10 μmol/L NGI-1 for 24 hours. p-eIF2α and actin were examined by immunoblotting. Scale bar, 100 μm. E, Representative paired IHC staining of B7-H4 and phospho-eIF2α (Ser51) in tissue array BC081120. Statistical analysis of IHC staining indicates B7-H4 expression is negatively correlated with p-eIF2α expression in breast cancer (r = −0.249, P = 8.71 × 10⁻³). F, MDA-MB-468-FLAG-hB7-H4 were treated in the presence or absence of doxorubicin (10 μmol/L) and/or NGI-1 (10 μmol/L). Then FLAG-hB7-H4 was immunoprecipitated followed by immunoblot. The indicated proteins were examined. G, Schematic diagram of the procedure of OptiPrep density gradient assay with 24 collected fractions from low to high density is shown. MDA-MB-468-vector and MDA-MB-468-B7-H4 knockout cells were treated with 10 μmol/L doxorubicin for 24 hours followed by OptiPrep density gradient assay. HSP90, CALR, eIF2α, and p-eIF2α in fraction 1 to 13 were examined by immunoblotting. PERK, eIF2α, and p-eIF2α were examined.

whether inhibition of DAMP exposure by B7-H4 would affect this pathway. Mouse B7-H4 expression in 4T1 cells was undetectable by immunoblot (Supplementary Fig. S6A). Real-time quantitative PCR showed that the Ct (cycle threshold) of mouse B7-H4 in 4T1 cells was more than 37 cycles, indicating no expression of mouse B7-H4 in 4T1 cells. Thus, mouse 4T1 breast cancer cells were stably transfected with vector or mouse B7-H4 (Supplementary Fig. S6A). The cells were then labeled with 5-(and-6)-(((4-chloromethyl) benzoyl)amino) tetramethylrhodamine (CMTMR), and treated with NGI-1 and/or doxorubicin, and then confronted with DCs isolated from the spleens of BALB/c mice.

The DCs were identified by immunostaining for CD11c. B7-H4 overexpression led to reduced phagocytosis (measured by the cooccurrence of CMTMR and CD11c) of the cancer cells by DCs, unless the cancer cells were treated with both NGI-1 and doxorubicin (Fig. 6A and B; Supplementary Fig. S6B and S6C). Of note, mB7-H4 overexpression was also linked to relatively reduced induction of the DC maturation markers CD86 and MHC class II (I-A/I-E; Supplementary Fig. S6D and S6E).

To determine the effects of B7-H4 in doxorubicin-induced ICD in vivo, we performed vaccination experiments in which immunocompetent BALB/c mice or T cell–deficient nu/nu
mice were first injected with cells cultured with doxorubicin and/or NGI-1 and then were challenged 1 week later with the same kind of live cells injected into the opposite mammary gland (Fig. 6C and D; Supplementary Fig. S6F). In nonvaccinated mice, 4T1-B7-H4 tumors grew faster than 4T1 vector alone in immunocompetent but not in immunodeficient hosts, 4T1-B7-H4 tumors grew transiently delayed in immunocompetent mice vaccinated with doxorubicin-treated cells, in line with the fact that 4T1 cells are poorly immunogenic. Intriguingly, combined doxorubicin/NGI-1 treatment yielded a more potent vaccine than doxorubicin treatment alone (Fig. 6C). In contrast, no differences were observed between the nonvaccinated and the vaccinated groups in nude mice (Fig. 6D), in line with the interpretation that T cells are required for the doxorubicin/NGI-1–elicited immunogenic effect.

IFNγ production by CD8+ cytotoxic T cells is essential for tumor growth control after activation of the immune response by cancer cells that succumb to ICD. The frequency of splenic CD8+ IFNγ+ T cells significantly decreased in mice bearing 4T1-B7-H4 tumors compared with mice with 4T1-vector tumors, as determined by immunofluorescence detection of intracellular IFNγ among CD8+ T lymphocytes. CD8+ IFNγ+ T cells were remarkably increased in mice vaccinated with doxorubicin + NGI-1–treated 4T1-B7-H4 cells than in mice vaccinated with doxorubicin only–treated cells (Supplementary Fig. S6G). The enzyme-linked immunospot (ELISPOT) assays confirmed these results at the level of splenic IFNγ production (Fig. 6E), and IHC at the level of CD8+ infiltration of the tumors (Supplementary Fig. S6H). In each case, doxorubicin-treated mB7-H4–overexpressing cancer cells induced a relatively poor CD8+ IFNγ+ T-cell immune response (compared with parental cancer cells expressing little or no CD8+) unless they were treated with both doxorubicin and the B7-H4 degradation–inducing drug NGI-1.

Modulation of B7–H4 Glycosylation and Ubiquitination Affects Tumor Growth and Doxorubicin-Induced Immunogenic Cell Death

4T1 breast cancer cells were engineered to stably express human (h) B7–H4 wild-type, ubiquitination-deficient hB7–H4–2KR mutant, or glycosylation-deficient hB7–H4–16NQ mutant (Supplementary Fig. S7A). No differences in cell growth and proliferation were observed in 4T1–hB7–H4, 4T1–hB7–H4–2KR, and 4T1–hB7–H4–16NQ cells (Supplementary Fig. S7B). The mutant B7–H4–16NQ protein (which cannot be glycosylated) was more restricted to the cytosol, while B7–H4 and B7–H4–2KR (which cannot be ubiquitinated) were localized in both the cytosol and the cell membrane (Supplementary Fig. S7C). B7–H4–16NQ was able to localize to the cell membrane, but at significantly lower levels than B7–H4–WT (Supplementary Fig. S7D and S7E), suggesting that the mutant retains partial functional activities of B7–H4 in terms of the membrane localization. Moreover, as compared with the unmutated hB7–H4, the glycosylation-deficient hB7–H4–16NQ mutant exhibited increased binding to AMFR, but lost binding to HSP90 (Supplementary Fig. S7F), indicating AMFR is crucial for the regulation of B7–H4–16NQ protein abundance.

In contrast, when inoculated into immunocompetent hosts, hB7–H4–2KR tumors progressed more quickly than 4T1–B7–H4 tumors, contrasting with the delayed growth of hB7–H4–16NQ tumors compared with 4T1–B7–H4 tumors. hB7–H4–WT and 16NQ tumors growth were significantly reduced in response to vaccination with cells treated with doxorubicin + NGI–1. In contrast, hB7–H4–2KR tumors did not respond to vaccination with cells treated with doxorubicin + NGI–1 (Fig. 6F; Supplementary Fig. S7G).

Splenic CD8+ IFNγ+ T cells were increased in mice bearing 4T1–hB7–H4–16NQ tumors, whereas CD8+ IFNγ+ T cells were decreased in 4T1–hB7–H4–2KR tumor bearers compared with those of the 4T1–B7–H4 group responding to the vaccination with doxorubicin + NGI–1. Similar results were obtained with ELISPOT assay (Fig. 6G and H; Supplementary Fig. S7H). Notably, we observed a 2-fold increase of tumor-infiltrating CD8+ T cells in 4T1–hB7–H4–16NQ tumors compared with that of the 4T1–B7–H4 group. No significant difference in CD8+ density between the vaccination group and nonvaccination group was observed for 4T1–hB7–H4–2KR tumors (Fig. 6I). Altogether, these results confirm that the abolition of B7–H4 glycosylation (hB7–H4–16NQ) and ubiquitination (hB7–H4–2KR) affected tumor cell immunogenicity in a positive and negative fashion, respectively.

Figure 6. B7–H4 and its ubiquitination or glycosylation-deficient mutants profoundly alter tumor growth and doxorubicin-induced immunogenic cell death. A and B, In vitro phagocytosis of coculture of mouse DCs and tumor cells. 4T1–vector and 4T1–B7–H4 cells were treated with doxorubicin (25 μM) or NGI–1 (10 μM) for 24 hours and cocultured with the purified CD11c+ cells for 2 hours at a ratio of 1:1, and then subjected to flow cytometry. n = 3 mice per group. C, In vivo vaccination assay. 4T1–vector or 4T1–B7–H4 cells were treated with doxorubicin (Dox; 25 μM) alone or in combination with NGI–1 (10 μM) for 24 hours. Then these cells (106 per mouse) were orthotopically injected into the right fourth mammary gland of the BALB/c mice (the vaccination step). PBS was used in the nonvaccinated group. One week later, all mice were rechallenged with live 4T1–vector or 4T1–B7–H4 cells (3 × 105 per mouse) of the same kind as the vaccination step in the left fourth mammary gland (the challenge step). The tumor growth was monitored twice per week. n = 8 mice per group. D, Failure of nu/nu BALB/c to mount an immune response against doxorubicin/NGI–1–treated 4T1–vector or 4T1–B7–H4 cells. Nude mice were inoculated with doxorubicin/NGI–1–treated 4T1–V or B7–H4 cells 1 to 2 weeks before the injection of live 4T1–vector or B7–H4 cells into the opposite flank; the tumor growth was monitored. n = 6 mice per group. E, 4T1–vector or 4T1–B7–H4 cells were treated with doxorubicin (25 μM) alone or in combination with NGI–1 (10 μM) for 24 hours. These cells were then orthotopically injected into the right fourth mammary gland of the BALB/c mice (the vaccination step). PBS was used in nonvaccinated control group. One or 2 weeks later, all mice were challenged with injection of live 4T1–vector or 4T1–B7–H4 cells (the same kind cells as used in the vaccination step) in the left fourth mammary gland (the challenge step). On day 28, mouse spleens of BALB/c mice were harvested and followed by ELISPOT. Quantification of IFNγELISPOT is shown. n = 3 mice per group. F–I, In vivo vaccination assays were performed with 4T1–hB7–H4, 4T1–hB7–H4–2KR, and 4T1–hB7–H4–16NQ cells. BALB/c mice were inoculated with doxorubicin/NGI–1–treated 4T1–hB7–H4, 4T1–hB7–H4–2KR, and 4T1–hB7–H4–16NQ cells 1 to 2 weeks (the vaccination step; PBS was used in the nonvaccinated group) before the injection of the live cells of the same kind (4T1–hB7–H4, 4T1–hB7–H4–2KR, and 4T1–hB7–H4–16NQ cells) into the opposite mammary gland (the challenge step); the tumor growth was monitored (F), n = 10 mice per group. G, On day 28, mouse spleens were harvested and followed by IFNγELISPOT. Quantification of IFNγELISPOT is shown. n = 3 mice per group. PBS is used in nonvaccinated group. H, On day 28, mouse spleens were harvested and followed by flow cytometry of staining IFNγ and CD8. Quantification of IFNγCD8–cells is shown. n = 3 mice per group. PBS is used in nonvaccinated control group. I, On day 28, mouse tumors in the mammary gland were harvested and digested followed by flow cytometry by detecting CD45 and CD8. n = 3 mice per group. PBS is used in nonvaccinated control group. Quantification of CD8–infiltrating cells is shown.
The Antitumor Effects of Combined Treatment with Doxorubicin and NGI-1 Are Exacerbated by PD-L1 Blockade

Previous in vivo vaccination experiments showed that combined doxorubicin/NGI-1 treatment yields a more potent vaccine than doxorubicin treatment alone (Fig. 6C), indicating that NGI-1 combined with doxorubicin could prevent tumor growth in settings of tumor cell vaccinations. Therefore, we investigated whether NGI-1 combined with immunogenic chemotherapy has antitumor efficacy against established tumors. The PD-1 pathway regulates activated T cells at the later stages of an immune response, primarily in peripheral tissues (39). B7-H4 is thought to inhibit T-cell responses at an early stage (40). We showed that NGI-1 inhibits B7-H4 significantly, and also inhibits glycosylation of PD-L1 (Supplementary Fig. S3D). However, NGI-1 also increased unglycosylation of PD-L1 (Supplementary Fig. S3D), indicating that the blocking of the PD-L1 signaling pathway might still be required. We therefore examined the effect of the triple combination of NGI-1, immunogenic chemotherapy, and anti–PD-L1 blocking antibody in mouse orthotopic tumor models of 4T1, where the basal mouse PD-L1 was knocked out followed by adding back human PD-L1 gene (Fig. 7A–D; Supplementary Fig. S8A–S8E) and E0771 (Fig. 7E–G; Supplementary S8F–S8I). Camsirubicin shares similar immunogenicity with doxorubicin, as we mentioned, and has been utilized in the triple combination.

Human PD-L1-expressing mouse tumor cells have been previously utilized to study therapeutic activity and mechanism of the action of anti-human PD-L1 in mice. The hPD-L1 mouse model system has facilitated the preclinical investigation of efficacy and immune-modulatory function of various forms of anti–hPD-L1 antibodies (41, 42). Given that intertumoral and intratumoral heterogeneity is a key challenge in cancer medicine (43), and based on our observation of an inverse correlation between B7-H4 and PD-L1 expression in cancer medicine (43), we showed that NGI-1 inhibits B7-H4 expression negatively correlated with that of PD-L1. The frequency of CD8+ T cells, in particular CD8+ IFNγ+ cells, was significantly increased by the combination of camsirubicin + NGI-1 + durvalumab compared with the monotherapies (Fig. 7C and D).

In addition, we generated E0771 cells expressing mouse B7-H4 expression for experimental use (Supplementary Fig. S8F). An E0771-mB7-H4 syngeneic C57BL/6 mouse model was established by mammary gland orthotopic injection of E0771-mB7-H4 cells, followed by injection of camsirubicin, NGI-1–releasing nanoparticles, and/or anti–mPD-L1 (clone:10F.9G2; Supplementary Fig. S8G). The administration of camsirubicin at 5 mg/kg, NGI-1 nanoparticles, and anti–mPD-L1 did not cause weight loss (Supplementary Fig. S8H). The combination of camsirubicin + NGI-1 + anti–mPD-L1 significantly decreased tumor growth (Fig. 7E) and caused the disappearance of more than 50% of E0771-mB7-H4 tumors (Fig. 7F). The frequency of CD8+ IFNγ+ T cells in spleens was significantly increased by the combination of camsirubicin + NGI-1 + anti–mPD-L1 compared with each monotherapy (Fig. 7G). Taken together, the above results suggest that the suppression of B7-H4 by NGI-1 may enhance the efficacy of PD-L1–targeted immunotherapy combined with immunogenic chemotherapy.

**DISCUSSION**

On the basis of unbiased bioinformatic analyses of immune-related proteins, we identified B7-H4 to be accumulated in immune-cold subtypes of TNBC tumors, and that B7-H4 expression negatively correlated with that of PD-L1. We also demonstrated that PTMs of B7-H4 dictate tumor immunogenicity in the context of doxorubicin-induced chemotherapy.
Regulation of B7-H4 by PTM Orchestrates Tumor Immunogenicity

**A**

![Graph showing tumor volume over days for 4T1-hB7-H4 and 4T1-hPD-L1 models](image)

**B**

![Graph showing percent survival over days for 4T1-hB7-H4 and 4T1-hPD-L1 models](image)

**C**

![Bar graph showing relative percentage (%) for different treatments](image)

**D**

![Image showing CD8 infiltrating cells in tumor (%) for different treatments](image)

**E**

![Graph showing tumor volume over days for E0771-mB7-H4 model](image)

**F**

![Graph showing tumor-free mice (%) for different treatments](image)

**G**

![Bar graph showing IFNγ/Cytotoxic T lymphocytes (%) for different treatments](image)

**H**

![Diagram showing cold tumor and hot tumor models](image)
immunogenic cell death. Pharmacologic inhibition of B7-H4 glycosylation by NGI-1 facilitated ubiquitination of B7-H4, thereby causing B7-H4 depletion and deactivation of ICD pathways, allowing the tumors to respond to a combination of ICD-eliciting chemotherapy and PD-L1 blockade (Fig. 7H). This study provides a new paradigm to develop strategies to treat patients with immune-cold TNBC.

The mechanisms rendering tumors immune-cold are manifold, including the lack of tumor antigen, defects in antigen presentation, absence of T-cell activation, and deficient homing of immune cells to the tumor bed (2). Chemotherapy-induced ICD converts malignant cells into vaccines and may improve T-cell priming, ultimately facilitating the T cell-mediated attack of residual cancer cells (13, 37). Other therapeutic approaches to transform cold tumors into hot ones include using oncolytic viruses, tumor vaccines, CD40 agonistic antibodies, cytokines, antiangiogenic therapy, adoptive T-cell therapy including chimeric antigen receptor (CAR) T cells, or T cell–recruiting bispecific antibodies (2).

B7-H4 is well demonstrated as a transmembrane protein that is expressed by tumor-infiltrating myeloid cells (44). However, the function of B7-H4 expressed by cancer cells is less understood. B7-H4 has been reported to be present in the plasma membrane (45), cytosol (22), and nucleus (46). We observed a strong intracellular localization of B7-H4 in tumor cells. Although the receptor of membrane B7-H4 remains unknown or controversial (47), it is generally recognized that cell surface–associated B7-H4 inhibits T-cell function (40). The function of the cytosolic pool of B7-H4 has been unclear. Our study suggests that cytosolic B7-H4 contained in malignant cells functions as a negative regulator of ICD. This effect apparently resides in the capacity of B7-H4 to inhibit the PERK-mediated phosphorylation of eIF2α that is required for the exposure of CARL at the cell surface as a “danger” or “eat-me” signal. Thus, B7-H4 subverts the entire cascade that usually allows ICD-associated DAMPs to elicit tumor cell phagocytosis by DCs, DC maturation, and T-cell recruitment to the tumor site. Fractionation analyses as well as co-IP experiments suggest that B7-H4 directly or indirectly interacts with eIF2α. This may either result in conformational changes of eIF2α that subsequently block its phosphorylation or sterically prevent eIF2α to bind to its kinases (such as PERK), a possibility that awaits further investigation.

Our studies uncovered a hidden B7-H4 protein turnover mechanism that can be ignited by the removal of N-linked glycosylation. B7-H4 is degraded through the ubiquitin–proteasome pathway, especially after the enzymatic removal or the pharmacologic or genetic inhibition of glycosylation. We found that the previously described glycosylation of B7-H4 (27) involves the hetero-oligomeric enzyme OST complex including its catalytic subunit STT3A. In addition, B7-H4 can be reglycosylated by UGCGG to facilitate its appropriate folding. We demonstrated that the glycosylation of B7-H4 within the ER ultimately determines the cellular abundance of the protein. Mass spectrometry identified five asparagine residues as N-linked glycosylation sites. However, the mutagenesis of these sites was not sufficient to generate an entirely glycosylation-deficient mutant, perhaps because additional asparagine residues could compensate for the function of the mutated sites. Thus, we generated unglycosylated B7-H4 mutant (B7-H4–16NQ). The mutant 16NQ is more flexible than B7-H4-WT in the Ig-like C2-type domain, though the whole structure is still stable by MD simulations. The 16NQ retains partial ability for trafficking to the membrane. Protein stability of 16NQ is independent from the HSP90-facilitated protein folding and degradation pathway; rather, it involves the AMFR-mediated proteasome–ubiquitin pathway, possibly because glycosylation deficiency of the 16NQ B7-H4 facilitates the binding with AMFR.

Several strategies have been developed to target B7-H4 in cancer treatment, including antibodies, CAR T cells, and drug conjugates. B7-H4 is reported to be an unstable cell-surface antigen, and hence may not be suitable as a target of antibody-dependent cellular cytotoxicity or mAbs (48). Here, we delineate an alternative therapeutic approach to target B7-H4 through glycosylation inhibition by NGI-1. NGI-1 suppresses the activity of OST through a direct and reversible interaction with both catalytic subunits, although it does not cause a complete disruption of glycosylation (as this would occur with tunicamycin), a fact that may explain its reduced toxicity as compared with other, more complete glycosylation inhibitors (31). NGI-1 is reported to block EGFR signaling in NSCLC and glioma (31–33). Our study suggests NGI-1 inhibits B7-H4 more potently than EGFR in all tested breast cancer cell lines. Whether NGI-1 efficiently targets other proteins awaits further investigation.

One critical challenge in cancer immunotherapy research is to develop more accurate preclinical models that may translate to humans. In our current work, to examine the antitumor effects of combined treatment with doxorubicin and NGI-1 in association with PD-L1 blockade, we have utilized a mouse orthotopic tumor model of 4T1-hPD-L1, where the basal mouse PD-L1 was knocked out followed by adding back human PD-L1 gene. Ideally, a mouse model that tests human immunologic genes in mice by knocking in the human gene could provide a more precise result. Coincidentally, 4T1 mouse breast cancer cells do not express B7-H4, which provides convenience to test the impact of B7-H4 in doxorubicin-induced immunogenic cell death in this work. Although the “humanization” of mouse cancer cells with human molecules is a possible strategy for studying the effects of therapeutic agents targeting such molecules, this system has some drawbacks resulting from incompatibilities between the two species, including immune responses to xenoaOntgens. Our results need to be further validated by more physiologically relevant models in the future translation study.

In summary, this study revealed a previously unidentified role for B7-H4 in determining local immunosuppression in the context of TNBC. B7-H4 protein stability is regulated by molecular cross-talk between ubiquitination and glycosylation, which subsequently dictates the efficacy of immunogenic (ICD-inducing) chemotherapy. Targeting B7-H4 for destruction by inhibiting its glycosylation, combined with immunogenic chemotherapy and PD-L1 blockade, could be an efficient strategy to treat TNBC, in particular when B7-H4 is abundantly expressed.
Regulation of B7-H4 by PTM Orchestrates Tumor Immunogenicity

STT3A (Bethyl Laboratories, A305–531A), and UGGG1 (Bethyl Laboratories, A302–026A), RPN2 (Bethyl Laboratories, A304–265A), Tech, 12034), mouse B7-H4 (R&D Systems, AF2154), RPN1 (Bethyl Signaling Technology, 5683), eIF2a (Sigma, HPA084885), AMFR (Cell Signaling Technology, 3452), rabbit IgG isotype control (Cell Signaling Technology, 3452), Alexa Fluor(R) 488 Conjugate (Cell Signaling Technology, 43405), anti-rabbit IgG (H+L), F(ab')2 Fragment (Alexa Fluor(R) 488 Conjugate; Cell Signaling Technology, 4414), rabbit IgG isotype control (Cell Signaling Technology, 3452), Alexa Fluor(R) 488 Conjugate (Cell Signaling Technology, 44125).

Cell Lines and Cell Culture

HEK293T, the authenticated breast cancer cell panel (#30–4500K), and the authenticated ovarian cancer cell panel (#TCP-1021), were obtained from ATCC in 2017. U2OS-HMGB1-GFP/H2B-RFP cells were provided by G. Kroemer (49) in 2018. 4T1-HDP-L1 cells, where the basal mouse PD-L1 was knocked out followed by adding back human PD-L1 gene, were provided by M.-C. Hung (41) in 2019. All cells were maintained in the media suggested by the ATCC such as 10% FBS, 100 U/mL streptomycin, and 100 U/mL penicillin. All cells were cultured at 37°C in a humidified atmosphere containing 5% CO2. All the cell lines were routinely tested for Mycoplasma infections. All cell lines were Mycoplasma tested every 3 months using MycoProbe Mycoplasma Detection Kit (R&D Systems). The length of time between cell line thawing and use in experiment does not exceed 1 month (two or more passages). All cell lines were authenticated by short tandem repeat DNA fingerprinting every 1 to 2 years and by visual inspection prior to experiments and freezing of cell line stocks in a central cell bank.

Chemicals and Antibodies

Camsirubicin was kindly provided by Monopar therapeutics Corporation. PEGase F was purchased from New England Biolabs Inc. Cycloheximide, doxorubicin, MG132, tunicamycin, Brefeldin A, puromycin dihydrochloride, and polyethyleneimine (PEI; branched from Polysciences, Inc.) were provided from MedChem Express. PEG (5000)-b-PLA (10,000; Diblock Polymer) was purchased from Sigma-Aldrich. NGI-1 and durvalumab were purchased from MedChem. DAB (DAKO) was used to visualize the reaction, followed by counterstaining with hematoxylin. After immunostaining, the sections were scanned at 20x magnification at standardized settings using a Nanozoomer (Hamamatsu) by a single investigator who was not informed of the clinical characteristics. The whole slide image was initially acquired in the NDPI File Format and further semiquantitatively analyzed by open-source software QuPath v0.2.0-m4 (50). The software was run on a standard off-the-shelf laptop computer. The H-score (or “histo” score) of B7-H4, CD8α, PD-L1, STT3A, AMFR, and phospho-eIF2α was assigned to tumor cells were evaluated by TMA algorithms. H-score is assigned using the following formula: H-score = 1x% of weakly staining cells + 2x% of moderately staining cells + 3x% of strongly staining cells + 4x% of weakly staining cells, giving a score range of 0–300.

Plasmids and Transfection

The N-FLAG tag human B7-H4 (HG10738-NA), N-MYC tag human B7-H4 (HG10738-NA), mouse B7-H4-C-FLAG tag (MGS0017-CF), and AMFR-C-MYC tag (MGS9687-CM) plasmids were ordered from Sino Biological Inc. pDNA-AMFR-C-FLAG (Plasmid #62370) and pCDNA-AMFR-C-FLAG RING mutant C356G H361A C-FLAG (Plasmid #61751; ref. 29) were purchased from Addgene. For plasmid transfection, cells were plated to form a 50% to 70% confluent culture. The HEK293T cells were transfected using Lipofectamine 2000 (Invitrogen). LentCas9-Blast (Plasmid #2962) was purchased from Addgene. All shRNA and CRISPR clones were purchased from Sigma Aldrich. For CRISPR B7-H4 knockout using sanger clone HSS000001377 and HSS000001378, lentCRISPRv2 puro (Addgene, Plasmid #98290) was used for the vector control. UGGG1 (SHCLNG-NM_19899, TRCN000110450 and SHCLNG-NM_198999, TRCN000110453), STT3A (SHCLNG-NM_008408, TRCN000093714 and SHCLNG-NM_008408, TRCN000093716), AMFR (SHCLNG-NM_001144, TRCN000003374 and SHCLNG-NM_001144, TRCN0000368947), RNPI (SHCLNG-NM_002950, TRCN0000421570), and RPN2 (SHCLNG-NM_002951, TRCN0000159696).
**Fragment Construction and Site-Directed Mutagenesis**

Indicated B7-H4 and AMFR fragments were amplified by PCR. All amplified products were further cloned into the pRES vector with FLAG-HA or HA tag. Fragment constructs were confirmed by DNA sequencing. B7-H4 2KR, 5NQ, 12NQ, 13NQ, and 16NQ mutations were introduced using mutagenic primers (QUICKCHANGE MULTISITE KIT, Agilent Technologies). Mutants were confirmed by sequence analysis.

**Lentiviral Construction, Generation, and Infection**

hB7-H4, hB7-H4–2KR, and hB7-H4–16NQ were generated by PCR amplification and were subcloned into pENTER-D-TOPO (Invitrogen) as an entry clone. In vitro recombination between an entry clone (containing a gene of interest flanked by attL sites) and a destination vector was performed to construct lentivirus expression vector. Clones with the right sequence were chosen.

Lentiviral particles are generated by transfection of the expression plasmid (i.e., pLenti-hB7-H4, hB7-H4–2KR and hB7-H4–16NQ), plus the packaged lentiviral particles (pSV-G, pRRE, and pRSV-REV) were collected, mixed with polybrene, and added into 293T cells using Lipofectamine 2000 (Life Technologies). Culture media were harvested 48 hours after transfection, filtered through 0.45-μm filters. 4T1 cells were transduced with the lentivirus. Upon infection, the stable cell lines were established by selecting with 2–5 μg/mL puromycin.

**Enzymatic Deglycosylation and Glycosylation Site Identification**

Deglycosylation experiments were performed on protein extracts from the indicated cells using Peptide N-Glycosidase F (New England Biolabs, Inc.) as per the manufacturer’s directions. Briefly, denaturing reaction conditions were used for PNGase F treatment, as follows. Cell lysates were denatured in PNGase F denaturing buffer (5% SDS and 10% 2-mercaptoethanol) for 10 minutes at 100°C. Denatured lysates were then cooled and mixed with 1/10 volume each of concentrated PNGase F reaction buffer, 0.5 mol/L sodium phosphate pH 7.5, and 10% NP-40. Samples were then digested with PNGase F for 1 hour at 37°C. The reaction was stopped by the addition of sample buffer containing 2-5 μg/mL puromycin.

**Purification of B7-H4 Complex**

MDA-MB-468 cells stable expression FLAG-B7-H4 were lysed with NP40 buffer (1% NP-40, 10% glycerol, 25 mmol/L Tris-HCl (pH 7.9) and protease inhibitor cocktail. B7-H4–interacting proteins were purified by immunoprecipitation on anti-FLAG M2 beads (catalog no. A2220, Sigma) and washed four times with TBST buffer (137 mmol/L NaCl, 20 mmol/L Tris-HCl (pH 7.6), 0.1% Tween-20). The complex was eluted with 3× FLAG peptide (Sigma, F4799) in TBS buffer. FLAG-B7-H4 protein was purified by immunopurification on anti-FLAG M2 beads (Sigma). The elute was then separated on SDS-PAGE followed by Coomassie blue staining. The bands were cut out for mass spectrum analysis. For the glycosylation sites, the elute was treated with Peptide N-Glycosidase F followed by SDS-PAGE. The bands were cut out at the bottom of stacking gel before running into the separating gel for mass spectrometry analysis.

**Cell Proliferation Assay**

Cells were seeded in 96-well plates at the concentration of 5,000 cells per well and incubated at 37°C for 3 days. After 3 days, the media were treated with antibodies diluted and 100 μL of phenol red-free DMEM with 10% CCK8 were added in each well and incubated for 1 hour. The absorbance of light with wavelength of 450 nm was measured with 800TS Absorbance Reader from the BioTek.

**Immunoblot Analysis**

Cells were lysed with 1 × cell lysis buffer (Cell Signaling Technology, 9806) containing protease inhibitor (Sigma-Aldrich, P8340) on ice for 10 minutes, homogenized by passing through a 21-gauge needle, and centrifuged at 14,000 × g for 15 minutes at 4°C to pellet the cell debris. Protein was quantified using the BCA assay (Thermo Fisher Scientific, 23225) and 20 μg of each sample was resolved on SDS-PAGE and transferred to nitrocellulose membrane. Membranes were blocked with TBST containing 5% skim milk for 1 hour and incubated overnight at 4°C with various primary antibodies followed by incubation with goat anti-rabbit or anti-mouse IgG horseradish peroxidase secondary antibody (Cell Signaling Technology, 7074 or 7076) at room temperature for 1 hour. Chemiluminescence substrate was applied using SuperSignal West Pico Chemiluminescent Substrate (Thermo Fisher Scientific, 34080) and blots were analyzed using the ChemiDoc Touch Imaging System (Bio-Rad). Semi-quantification of data was performed using Image Lab (Bio-Rad).

**Immunoprecipitation Analysis**

Cells were lysed at 4°C in ice-cold immunoprecipitation assay buffer (Cell Signaling Technology, 9806) and cell lysates were cleared by brief centrifugation (13,000 × g, 15 minutes). Concentrations of proteins in the supernatant were determined using the BCA assay. For immunoprecipitation assay, cell lysate was incubated with anti-FLAG M2 beads or IP-proved antibodies overnight at 4°C on a rotator, followed by the addition of protein A/G plus agarose to the reaction for 2 hours at 4°C. To detect ubiquitin conjugates, the lysates were denatured in lysis buffer containing 1% SDS to disrupt any protein-protein interactions, and then diluted approximately 10-fold prior to IP followed by immunoblot. Prior to immunoprecipitation, samples containing equal amounts of proteins were precleared with protein A agarose beads (Santa Cruz Biotechnology, sc-2027; 4°C, 3 hours), and subsequently incubated with various irrelevant IgG or specific antibodies in the presence of protein A agarose beads for 2 hours or overnight at 4°C with gentle shaking. After five washes with lysis buffer supplemented with protease inhibitor mixture, complexes were released from the anti-FLAG M2 beads by boiling for 5 minutes in 2× SDS-PAGE loading buffer.

**Flow Cytometry**

Single-cell suspensions were stained with relevant mAbs and then washed twice with cold PBS. Intracellular IFNγ staining under stimulation with 10 ng/mL PMA, 100 ng/mL ionomycin plus 10 μg/mL BFA for 4 hours was performed according to the manufacturer’s instructions (BD Biosciences) and were stained with surface markers and DEAD Fixable Violet Dead Cell Stain Kit (Thermo Fisher Scientific, # L34963) before fixation in 4% formaldehyde for 10 minutes at room temperature. Cells were washed with ice-cold PBS containing 2% BSA, followed by the second wash step with ice-cold PBS. Cells were resuspended in 80% methanol and incubated for 30 minutes at −20°C. The pellet was washed twice with ice-cold PBS. Annexin V Alexa Fluor 647 Ready Flow Conjugate (Thermo Fisher Scientific, #R37175) and DAPI staining were used to measure markers and DEAD Fixable Violet Dead Cell Stain Kit (Thermo Fisher Scientific, # L34963) before fixation in 4% formaldehyde for 10 minutes at room temperature. Cells were washed with ice-cold PBS containing 2% BSA and analyzed by flow cytometry. Samples were conducted on a BD FACSymphony A5-Laser Analyzer, and data were analyzed with FlowJo software.

**Phagocytosis and DC Maturation Assay**

The cells isolated from spleen of three naive mice were harvested, followed by the purification of CD11c+ cells using CD11c Microbeads UltraPure (Miltenyi Biotec, # 130–108–338). Tumor cells were stained with CellTracker Orange CMTMR (Thermo Fisher Scientific, #C2927) at 37°C for 30 minutes, and then treated without
doxorubicin (25 μmol/L) or NGI-1 (10 μmol/L) for 24 hours. These tumor cells were cocultured with the CD11c+ cells for 2 hours at a ratio of 1:1, and then subjected to flow cytometry (phagocytosis assay). For dendritic maturation assay, tumor cells were cocultured with the CD11c+ cells for 2 hours at a ratio of 1:1, and then subjected to flow cytometry (13).

**OptiPrep Density Gradient Protein Fractionation Assay**

To prepare the iodixanol gradient, 50% (v/v) and 5% (v/v) solutions of iodixanol were made by diluting the stock solution (60% (v/v) aqueous iodixanol from StemCell Technologies) with 0.25 mol/L sucrose, 4 mmol/L EDTA, 60 mmol/L Tris-HCl pH7.4. The gradient was formed by gradient makers and loaded into a 14 × 89 mm polycarbonate tube (Beckman Coulter). Cells were lysed with 0.5% NP40, and 5 mg protein was overlaid onto the top of the gradient, and centrifugation performed at 100,000 × g for 18 hours at 4°C. Twenty-four individual 0.5-mL gradient fractions were collected with increasing density.

**Immunofluorescence**

Cells grown on coverslip were fixed with 4% paraformaldehyde in PBS for 10 minutes. The membrane staining of the indicated proteins was performed without membrane permeabilization by 0.1% Triton X-100 unless otherwise stated. The indicated antibodies were incubated with cells 4°C overnight. After three washes with PBSB, cells were incubated with Alexa 488 anti-rabbit/594-anti-mouse IgG diluted in PBSB for 1 hour. Cells were washed, mounted with UltraCruz DAPI containing mounting medium (Santa Cruz Biotechnology), viewed, and photographed under Nikon A1 (Nikon) with the software of NIS-Elements. The fluorescence of doxorubicin alone was also checked.

**Duolink Proximity Ligation Assay**

The colocalization of B7-H4, AMFR, STT3A, and CALR was analyzed by the proximity ligation assay by using Duolink In Situ PLA Kit (Sigma-Aldrich) and following the manufacturer’s protocol. In brief, cells were plated on glass coverslips. Then the cells were fixed with 4% paraformaldehyde followed by permeabilization (0.1% Triton X-100). Cells were then incubated with blocking buffer followed by primary antibody overnight at 4°C. Cells were then incubated with PLA probes. Ligation mix was added to cells for 1 hour at 37°C. The amplification mix was added for 2 hours at 37°C and the cells were washed twice. The coverslips were mounted using the Duolink In Situ Mounting Medium followed by imaging the cells with BioTek LionHeart automated fluorescence microscope.

**Structure Modeling and Docking Analysis**

Structural models for B7-H4 Ig-like V-type 1 and V-type 2 domains were generated by comparative/homology modeling, searching the Swiss-Model server. Protein–protein docking simulations for AMFR/RING domain (PDB: 2LXH) and the modeled B7-H4 structure were performed using the ZDOCK server. Docking poses were ranked on the basis of the ZDOCK 3.0.2’s scoring function and the top 100 binding poses were clustered using K-means clustering with a cutoff RMSD of 5 Å into five clusters. Docking poses with the highest scores in each cluster were selected to conduct two independent 120-ns unbiased MD simulations to further evaluate the stability of the predicted binding poses. NAMD package and CHARMM36m force field were used in MD simulations.

**In Vivo Vaccination Assay**

BALB/c and nu/nu mice (6-week-old female) were purchased from Charles River. Stable expression of empty-vector 4T1 (4T1-vector) or 4T1-B7-H4 wild-type of 4T1 (4T1-B7-H4) cells were treated with doxorubicin (25 μmol/L) alone or in combination with NGI-1 (10 μmol/L) for 24 hours. Then, the cells were collected and orthotopically injected (10⁶ cells per mouse) into the right fourth mammary gland of immunocompetent BALB/c mice. PBS was used in nonvaccinated control group. One week later, all mice were rechallenged with the same kind of live 4T1-vector or 4T1-B7-H4 cells (3 × 10⁶ per mouse) in the left fourth mammary gland. The tumor growth was monitored twice per week. All animal experiments were approved by the Institutional Animal Care and Use Committee (IACUC) of Northwestern University (Chicago, IL).

**In Vivo Therapeutic Assay**

NGI-1 nanoparticles were prepared as reported previously (33). Briefly, polyethylene glycol (PEG)-b-Polyactic acid nanoparticles were synthesized using diblock polymer (Polysciences, #25018) and Polyethyleneimine (Sigma-Aldrich, #408719) using a nanoprecipitation technique. Polymer was dissolved in DMSO and was added drop-wise to deionized water to create an NP suspension followed by Amicon Ultracell 100k centrifugal filter unit, and centrifuged at 4,000 × g at 4°C for 30 minutes. For drug-loaded NPs, NGI-1 was dissolved in DMSO at a concentration of 50 mg/mL, and PEI was then dissolved in DMSO at 50 mg/mL. The NGI-1 and PEI solutions were then mixed at a 6:1 ratio (by weight) of PEI:drug and then added to the PLA-PEG solution at a 10% ratio of NGI-1:PLA-25 PEG by weight. A mixture of 4T1-hPD-L1 (2 × 10⁶; ref. 41) and 4T1-B7-H4 (2 × 10⁵) cells at a ratio of 1:1 or E0771-mB7-H4 (1 × 10³) was orthotopically injected into the left fourth mammary fat pad followed by injection of camsirubicin intraperitoneally at dose of 25 mg/kg or 5 mg/kg for a total of four times. The NGI-1 nanoparticles was then injected intravenously at 10 mg/kg four times, followed by the administration of PD-L1 antibody durvalumab or inVivoMAB anti-mPD-L1(10F.9G; BioXCell, catalog no.: BE0101) intraperitoneally at 5 mg/kg for a total of three times. The tumor growth was monitored twice per week. All animal experiments were approved by the IACUC of Northwestern University.

**ELISPOT Assay**

For detection of IFNγ-secreting cells, spleens were harvested at day 20 after tumor cell injection and stimulated with 10 ng/mL PMA, 100 ng/mL ionomycin ex vivo with 2 × 10⁵ cells/well for 24 hours. ELISPOT assay was performed with Mouse IFNγ ELISPot Kit (Thermo Fisher Scientific, #EL485) according to the manufacturer’s instructions. The numbers and diameters of spots were counted in duplicate, and calculated by an automatic ELISPOT counter (CTL ImmunoSpot S5 Macro Analyzer, Cellular Technology Limited).
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**Authors’ Contributions**

**X. Song:** Conceptualization, software, formal analysis, supervision, investigation, methodology, writing-original draft, writing-review and editing. **Z. Zhou:** Conceptualization, supervision, investigation, writing-original draft, project administration, writing-review and editing. **H. Li:** Software, formal analysis, investigation, methodology. **Y. Xue:** Resources, software, formal analysis, investigation. **X. Lu:** Resources, writing-review and editing. **I. Bahar:** Resources, software, writing-review and editing. **O. Kepp:** Resources, writing-review and editing. **M.-C. Hung:** Resources, writing-review and editing. **G. Kroemer:** Resources, writing-review and editing. **Y. Wan:** Conceptualization, data curation, supervision, funding acquisition, project administration, writing-review and editing.

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**REFERENCES**

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