

1 *Review*

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3 **HER2 directed antibody-drug-conjugates beyond** 4 **T-DM1 in breast cancer**

5 **Gabriel Rinnerthaler**^{1,2}, **Simon Peter Gampenrieder**^{1,2} and **Richard Greil**^{1,2,*}

6 ¹ Department of Internal Medicine III with Haematology, Medical Oncology, Haemostaseology, Infectiology
7 and Rheumatology, Oncologic Center, Salzburg Cancer Research Institute - Laboratory for Immunological
8 and Molecular Cancer Research (SCRI-LIMCR), Paracelsus Medical University Salzburg, Salzburg, Austria;
9 E Mails: g.rinnerthaler@salk.at (G.R.); s.gampenrieder@salk.at (SP.G.); r.greil@salk.at (R.G.)

10 ² Cancer Cluster Salzburg, Salzburg, Austria

11 * Correspondence: E-Mail: r.greil@salk.at Tel.: +43 5 7255 25801; Fax: +43 5 7255 25999.

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14 **Abstract:** Since the discovery of the human epidermal growth factor receptor 2 (HER2) as an
15 oncogenic driver in a subset of breast cancers and the development of HER2 directed therapies, the
16 prognosis of HER2 amplified breast cancers has increased meaningfully. Next to monoclonal anti-
17 HER2 antibodies and tyrosine kinase inhibitors, the antibody-drug conjugate T-DM1 is a pillar of
18 targeted treatment of advanced HER2-positive breast cancers. Currently, several HER2 directed
19 antibody-drug conjugates are under clinical investigation for HER2 amplified but also HER2
20 expressing but not amplified breast tumors. In this article, we review the current preclinical and
21 clinical evidence of the investigational drugs A166, ALT-P7, ARX788, DHES0815A, DS-8201a, RC48,
22 SYD985, MEDI4276 and XMT-1522.

23 **Keywords:** ADC; HM2-MMAE; (vic-)trastuzumab duocarmazine; Trastuzumab deruxtecan; TAK-
24 522; Trastuzumab emtansine; anti-HER2/PBD-MA; HER2 low; HER2-low; mode of action

25

26 **1. Introduction**

27 The human epidermal growth factor receptor 2 (HER2), known as erbB-2, or proto-oncogene Neu, is
28 a receptor tyrosine-protein kinase encoded by the ERBB2 gene on chromosome 17q12. Besides
29 epidermal growth factor receptor (EGFR, ErbB-1), human epidermal growth factor receptor 3 (HER3,
30 ErbB-3), and human epidermal growth factor receptor 4 (HER4, ErbB-4), HER2 is a member of the
31 epidermal growth factor (EGF) receptor family. Since the HER2 protein has no ligand binding
32 extracellular domain, no growth factors can directly bind to it. However, it forms heterodimers with
33 ligand-binding members of the EGF receptor family, stabilizing ligand binding and enhancing
34 kinase-mediated downstream signaling, including activation of phosphatidylinositol-3 kinase and
35 mitogen-activated protein kinase [1,2].

36 HER2 expression can be detected on cell membranes of epithelial cells in the gastro-intestinal tract,
37 respiratory tract, reproductive tract, urinary tract, skin, breast and placenta, but also on heart and
38 skeletal muscle cells [3] [4]. In fetal tissue, the level of HER2 expression is generally higher than in
39 corresponding normal adult tissue [4].

40 A HER2 amplification can promote tumorigenesis through multiple mechanisms and can therefore
41 be considered as an oncogenic driver in HER2 amplified cancers [1]. Despite breast cancer, HER2 was
42 found to be amplified and/or overexpressed in several cancer types including gastric and lung cancer
43 [5].
44

45 Approximately 15% of all breast cancer cases belong to the HER2-positive subtype defined by HER2
46 protein overexpression and/or HER2 gene amplification [6]. Traditionally, HER2-positive breast
47 cancer is regarded as the most aggressive subtype and a high rate of recurrences were observed before
48 the introduction of anti-HER2 targeted therapies. The addition of trastuzumab, a humanized
49 monoclonal antibody targeting HER2, to conventional adjuvant chemotherapy, however, resulted in
50 a significant and clinically relevant reduction of disease free survival (HR 0.60; 95% confidence
51 interval [CI] 0.50 - 0.71, $P < 0.001$) and overall survival (HR 0.66; 95% CI 0.57 - 0.77, $P < 0.00001$) [7].
52 Besides trastuzumab, further HER2-directed drugs such as the monoclonal antibody pertuzumab,
53 the antibody-drug conjugate (ADC) trastuzumab-emtansine (T-DM1) and tyrosine-kinase inhibitors
54 such as lapatinib and neratinib are available today, allowing targeted combination therapy or
55 sequential administration of non-cross resistant drugs [8].
56

57 In about 50% of breast cancers a low-level expression of HER2 without HER2 amplification can be
58 observed [9,10]. In two landmark adjuvant trastuzumab trials including patients with HER2-
59 amplified or overexpressing breast cancer according to local site laboratories, a cohort of patients
60 without HER2-amplification nor HER2 overexpressing by central testing was identified. These HER2-
61 low cohorts seemed to benefit from trastuzumab in a retrospective unplanned subgroup analysis
62 [11,12]. The efficacy of an adjuvant trastuzumab treatment in HER2-low (immunohistochemistry
63 [IHC] 1+ or 2+ but not HER2 amplified) breast cancer patients was prospectively investigated in the
64 phase III trial NSABP B-47 [13]. In this trial, 3,270 patients were randomized 1:1 to standard adjuvant
65 chemotherapy with or without one year of trastuzumab. No difference in regard to the 5-year disease-
66 free survival (DFS) was observed between the treatment groups. The findings did not differ according
67 by HER2 IHC level, extent of lymph node involvement, or hormone receptor status [13]. Despite
68 HER2 amplification as a predictor for trastuzumab benefit, we could recently demonstrate that a
69 poly-ligand profiling can differentiate trastuzumab-treated breast cancer patients according to their
70 outcomes [14].
71

72 Antibody-drug conjugates (ADCs) are molecules consisting of a recombinant monoclonal antibody
73 covalently bound to a cytotoxic drug (called drug payload or warheads) via a synthetic linker [15].
74 ADCs combine the advantage of antibodies in binding a specific target and the cytotoxic capability
75 of a chemotherapeutic drug. A stable linker between the antibody and the cytotoxic drug is crucial
76 for the ADC integrity in circulation. After antibody binding to the specific antigen on the (cancer) cell
77 surface, the ADC gets internalized and the cytotoxic drug is released intracellularly where it can exert
78 its effect. Using cleavable linkers, ADCs can be designed to promote drug release from the target cell
79 to the extracellular space. Thereby, surrounding and bystander cells, which may or may not express
80 the ADC target antigen, can be killed by taking up the cytotoxic drug [15,16]. This bystander killing
81 can also occur if the cytotoxic drug is released from the antibody after antigen binding just before
82 internalization. The supposed mode of action of ADCs in HER2-low breast cancer patients is outlined
83 in Figure 1.
84

85 T-DM1 is at present the only approved ADC for treatment of advanced HER2-positive breast cancer,
86 based on the phase 3 registration trials EMILIA [17] and THERESA [18] comparing T-DM1 with
87 capecitabine plus lapatinib and treatment of physicians choice, respectively. Recently, results of the
88 phase 3 trial KATHRINE, where adjuvant T-DM1 was compared to trastuzumab in HER2-positive
89 patients with residual disease after neoadjuvant chemo and anti-HER2 treatment, were published
90 [19]. Because of the favorable efficacy of T-DM1, an approval in the post-neoadjuvant setting is
91 awaited already in 2019.
92

93 T-DM1 is a second-generation ADC consisting of the monoclonal HER2 directed IgG1 antibody
94 trastuzumab, a non-cleavable thioether linker attached to random lysins and 3 to 4 maytansinoid
95 emtansine, also called DM1 [14]. DM1 has in vitro a 11 to 25X higher cytotoxic potency than

96 maytansine and is 24- to 270X more effective than taxanes. The mean ratio of DM1 molecules per
 97 antibody (drug antibody ratio, DAR) is 3.5 [20].

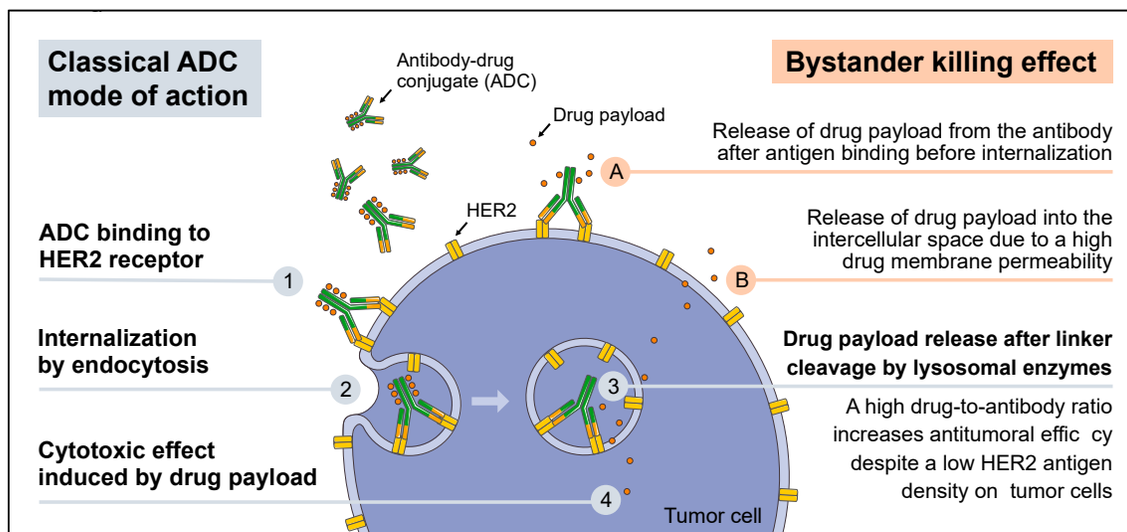
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100 Currently, several ADCs are under clinical investigation for breast cancer treatment. Most of the
 101 drugs target HER2, but also other receptors like HER3, the zinc transporter LIV1, receptor tyrosine
 102 kinase-like orphan receptor 2 (ROR2) and Trop-2 serve as targets for the investigational drugs (Table
 103 1 and Table 2). In this article we review the current evidence of investigational HER2 directed ADCs
 104 for the treatment of breast cancer.

104

105 **Figure 1. Mode of action of HER2 directed ADCs in HER2-low tumors**



106

107 **Legend.** Classical mode of action of ADCs with cleavable linkers: (1) After binding of the monoclonal
 108 anti-HER antibody component to HER2 expressed on the cell surface of tumor cells, (2) the ADC-
 109 HER2 complex is internalized by endocytosis. (3) After linker cleavage by lysosomal proteases, the
 110 drug payload is released and (4) can induce the cytotoxic effect leading to tumor cell death. A high
 111 drug-to-antibody ratio can increase antitumoral efficacy despite a low HER2 antigen density on
 112 tumor cells.

113 Bystander killing effect: Using cleavable linkers, ADCs can be designed to promote drug release from
 114 the target cell to the extracellular space. Thereby, surrounding and bystander cells, which may or
 115 may not express the ADC target antigen, can be killed by taking up the cytotoxic drug. (A) This
 116 bystander killing can occur if the cytotoxic drug is released from the antibody after antigen binding
 117 before internalization. (B) Additionally, the drug payload can be released from the tumor cell into the
 118 intracellular space due to a high membrane-permeability of the ADC drug payload. This figure was
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121

122 **Table 1: Investigational HER2 targeting antibody drug conjugates in breast cancer**

Drug name	Cytotoxic payload	Reported efficacy in HER2-low	Phases (number of trials, NCT identifier)	Company
A166	NA	no	Phase I/II: 1 (NCT03602079)	Klus Pharma Inc.

ALT-P7 (HM2-MMAE)	monomethyl auristatin E	no	I: 1 (NCT03281824)	Alteogen, Inc.
ARX788	monomethyl auristatin F	no	I: 2 (NCT02512237, NCT03255070)	Ambrx, Inc.
DHES0815A (anti-HER2/PBD-MA)	PBD-MA	no	I: 1 (NCT03451162)	Genentech, Inc.
DS-8201a (Trastuzumab deruxtecan)	DXd	yes	I: 3 (NCT03523572, NCT03368196, NCT03366428) II: 1 (NCT03248492) III: 3 (NCT03734029, NCT03523585, NCT03529110)	Daiichi Sankyo, Inc.
MEDI4276	AZ13599185	yes	-	MedImmune LLC
RC48	monomethyl auristatin E	no	Ib/II: 1 (NCT03052634) II: 1 (NCT03500380)	RemeGen
SYD985 ([vic-]trastuzumab duocarmazine)	seco-DUBA	yes	III: 1 (NCT03262935)	Synthon Biopharmaceuticals BV
T-DM1 (Trastuzumab emtansine)	DM1	no	I: 3 (NCT02073916, NCT02038010, NCT03364348) II: 3 (NCT03587740, NCT02073487, NCT02414646)	Roche
XMT-1522 (TAK-522)	AF-HPA	yes	I: 1 (NCT02952729)	Mersana Therapeutics

123 (clinicaltrials.gov, last access on 20th of December 2018)

124 NA: not available; PBD-MA: pyrrolo[2,1- c][1,4]benzodiazepine monoamide; seco-DUBA: synthetic
125 duocarmycin analogon seco-DUocarmycin-hydroxyBenzamide-Azaindole; AF-HPA: Auristatin F-
126 hydroxypropylamide

127

128 **Table 2: Investigated ADCs in breast cancer targeting receptors other than HER2**

Drug	Target	Running trials (number of trials, NCT identifier)	Company
U3-1402	HER3	Phase I/II: 1 (NCT02980341)	Daiichi Sankyo, Inc.
SGN-LIV1A	LIV1	Phase I: 1 (NCT01969643)	Seattle Genetics, Inc.
CAB-ROR2-ADC	ROR2	Phase I/II: 1 (NCT03504488)	BioAtla, LLC
Sacituzumab govitecan (IMMU-132)	Trop-2	Phase I/II: 1 (NCT01631552) Phase II: 1 (NCT02161679)	Immunomedics, Inc.

129 (clinicaltrials.gov, last access on 20th of December 2018)

130 ROR2: Receptor tyrosine kinase-like orphan receptor 2

131

132 **2. A166**

133 **Drug structure.** The ADC A166 is composed of a monoclonal anti-HER2 antibody conjugated to a
134 cytotoxic agent. Both, the monoclonal antibody and the cytotoxic agent are as of yet undisclosed [21].

135 **Ongoing trials without published results.** A166 is currently investigated in a running phase 1/2 trial
136 including patients with relapsed or refractory HER2 expressing or HER2 amplified cancers including
137 breast cancer patients (clinicaltrials.gov identifier: NCT03602079). After defining the maximum
138 tolerated dose (MTD) in the phase 1 dose escalation part of the trial, patients will be enrolled into
139 several cohorts including HER2 positive breast cancer patients (cohort 1) and HER2-low breast cancer
140 patients (IHC 1+ or 2+ but not HER2 amplified; escalation cohort 3).

141 To the best of our knowledge, no published data of A166 are currently available.

142

143 **3. ALT-P7 (HM2/MMAE)**

144 **Drug structure.** ALT-P7 is an ADC composed of the trastuzumab biobetter HM2 conjugated in a site-
145 specific manner to monomethyl auristatin E (MMAE) [22]. MMAE is a cytotoxic agent acting by
146 inhibiting the tubulin polymerization in dividing cells resulting in an in G2/M phase arrest and
147 apoptosis [23].

148 **Ongoing trials without published results.** ALT-P7 is currently investigated in an ongoing open-
149 label, dose escalation and phase 1 trial in patients with HER2 positive MBC, who have progressed on
150 previous trastuzumab-based therapy (clinicaltrials.gov identifier: NCT03281824).

151 To our knowledge, no published data of ALT-P7 are currently available.

152

153 **4. ARX-788 (ARX788)**

154 **Drug structure.** The novel ADC ARX-788 is composed of a monoclonal HER2 targeting antibody site-
155 specifically conjugated, via a non-natural amino acid linker para-acetyl-phenylalanine (pAcF), to
156 monomethyl auristatin F (MMAF) [24]. The site-specific conjugation of MMAF to the HER2 antibody
157 improves the therapeutic window of ARX-788 by increasing payload stability and optimizing its half-
158 life. The mean DAR is 1.9

159 **Preclinical data.** In murine xenograft models of the HER2-positive breast cancer cell lines BT474 and
160 HCC1954, a rapid tumor regression was induced after a single injection of ARX-788. In a
161 trastuzumab-resistant breast cancer xenograft model (JIMT-1), ARX-788 was significantly more
162 effective than T-DM1 in inducing tumor regression. A long-term stability of 12 and 8 days was
163 revealed in rodent and non-human primates, respectively. The conjugated form of ARX-788
164 remained intact over the course of a 3-week study in non-human primates.

165 **Ongoing trials without published results.** ARX-788 is currently investigated in two ongoing phase
166 I trials. The first part of both trials (phase 1a) is designed to determine the recommended phase 2 dose
167 (RP2D) in patients with HER2 positive advanced solid tumors. In the second part (phase 1b) of the
168 first trial, safety and activity of the RP2D will be tested in three expansion cohorts: a HER2-positive
169 advanced breast cancer cohort, a HER2-low advanced breast cancer cohort, and a HER2 positive
170 gastric cancer cohort (clinicaltrials.gov identifier: NCT02512237). The phase 1b of the second trial is
171 designed to investigate the activity and safety of the RP2D in two advanced breast cancer expansion

172 cohorts: one cohort with HER2 positive patients, and one cohort with HER2-low patients
173 (clinicaltrials.gov identifier: NCT03255070).

174

175 **5. DHES0815A (anti-HER2/PBD-MA)**

176 **Drug structure.** DHES0815A consist of a monoclonal HER-2 targeting antibody linked to pyrrolo[2,1-
177 c][1,4]benzodiazepine monoamide (PBD-MA) [25]. PBD-MA crosslinks DNA minor grooves, leading
178 to DNA strand breaks, cell cycle arrest, and cell death.

179 **Ongoing trials without published results.** DHES0815A is currently investigated in a first-in-human,
180 open-label, multicenter, dose-escalation phase 1 trial evaluating the safety, tolerability, and
181 pharmacokinetics (clinicaltrials.gov identifier: NCT03451162).

182 To our knowledge, no published data of DHES0815A are currently available.

183

184 **6. DS-8201a (Trastuzumab deruxtecan)**

185 **Drug structure.** DS-8201a is a novel ADCs composed of trastuzumab, an enzymatically cleavable
186 maleimide glycynglycyn-phenylalanyn-glycyn (GGFG) peptide linker and a topoisomerase I
187 inhibitor [26]. Topoisomerase I inhibitors induce double-strand DNA breaks and apoptosis by
188 binding to and stabilization of topoisomerase I-DNA cleavable complexes [27]. DXd, the
189 topoisomerase I inhibitor component of DS-8201a, is a derivative of exatecan mesylate (DX-8951f). In
190 various tumor xenograft models, including CPT-11-resistant tumors, antitumor activity of DX-895
191 was superior to irinotecan (CPT-11) [28]. Each trastuzumab molecule of DS-8201a is conjugated with
192 8 molecules of DXd. This DAR of 8 is higher compared to T-DM1 with a DAR of 3-4. After binding to
193 HER2 on the cell surface, DS-8201a gets internalized and the linker is cleaved by lysosomal enzymes
194 such as cathepsins B and L which are highly expressed in tumor cells.

195 **Preclinical data.** A potent bystander effect of DS-8201a is suggested due to a high membrane-
196 permeability of the DS-8201a payload DXd [29]. In comparison, the payload of T-DM1, Lys-SMCC-
197 DM1, has a low level of permeability. In coculture experiments of HER2-positive KPL-4 cells and
198 HER2-negative MDA-MB-468 cells, DS-8201a killed both cells, whereas T-DM1 could not. This
199 observation was confirmed in a xenograft model [15]

200 In various mice xenograft models with different HER2 expression levels, DS-8201a was also effective
201 in HER2 moderate positive and HER2 weak positive models, while T-DM1 was only effective in the
202 HER2 strong positive model [29]. In HER2-low patient derived xenografts (PDX), an antitumor
203 activity of DS-8201a but not of T-DM1 could also be shown.

204 **Clinical data.** Single agent DS-8201a is currently investigated in a large phase 1 trial in heavily
205 pretreated patients with HER2 expressing solid tumors, including patients with breast cancer
206 (clinicaltrials.gov identifier NCT02564900) [30,31]. Twenty-four patients were enrolled into the dose
207 escalation part (part 1), and further 260 patients are planned to be enrolled into several dose
208 expansion cohorts (part 2). In part 1, DS-8201a was administered up to 8.0 mg/kg. No dose limiting
209 toxicity was observed and the maximum tolerated dose (MTD) was not reached. For part 2, dose
210 levels of 6.4 and 5.4 mg/kg IV every 3 weeks were chosen.

211 In 99% of patients treated with 5.4 or 6.4. mg/kg (N=241, data cutoff April 2018), an adverse event
212 (AE) of any grade was observed. AEs \geq grade 3 occurred in 42% of patients and serious adverse events
213 were reported in 21% of patients. The most common non-hematological AEs were nausea (all grades:

214 69%, grade ≥ 3 : 3%), vomiting (all grades: 35%, grade ≥ 3 : 2%), diarrhea (all grades: 27%, grade ≥ 3 :
215 1%), decreased appetite (all grades: 56%, grade ≥ 3 : 3%), alopecia (all grades: 36%) and fatigue (all
216 grades: 28%, grade ≥ 3 : 2%). Anemia (all grades: 32%, grade ≥ 3 : 15%), thrombocytopenia (all grades:
217 29%, grade ≥ 3 : 10%) and neutropenia (all grades: 25%, grade ≥ 3 : 15%) were common. The frequency
218 of infusion-related reactions (all grades: 2%, grade ≥ 3 : 0%) was low. Laboratory abnormalities of liver
219 enzymes was generally of low grade (AST increase: all grades: 20%, grade ≥ 3 : 1%; ALT increase: all
220 grades: 16%, grade ≥ 3 : 1%). A decrease in ejection fraction (all grades: 1%, grade ≥ 3 : 0%) and a QT
221 prolongation (all grades: 5%, grade ≥ 3 : <1%) were uncommon. Interstitial lung disease (ILD; all
222 grades: 3%, grade ≥ 3 : 1%) and pneumonitis (all grades: 7%, grade ≥ 3 : 2%) were infrequent, but 5
223 fatal cases of ILD and pneumonitis were observed.

224 As of April 2018, 111 patients with HER2-positive metastatic breast cancer evaluable for efficacy
225 outcome were enrolled in this phase 1 trial, with a median age of 55 (range 33-77) and a median of 7
226 prior therapies (range 2-21) [31]. The confirmed overall response rate (ORR) was 55% with a disease
227 control rate (DCR: CR + PR + SD) of 94%. Median duration of response and median progression-free
228 survival (PFS) were not reached. Confirmed ORRs at dose levels 5.4 and 6.4 mg/kg were 53% and
229 56%, respectively [32]. The pharmacokinetic relationship between minimum blood plasma
230 concentration C_{min} of intact DS-8201a and ORR was statistically significant ($P=0.035$). Based on logistic
231 regression, a statistically significant relationship was observed between exposures and the following
232 AEs: neutropenia (any grade, $P=0.003$; grade ≥ 3 , $P=0.037$), anemia (any grade, $P=0.002$; grade ≥ 3 ,
233 $P<0.001$), thrombocytopenia (any grade, $P=0.021$), ILD/pneumonitis (any grade, $P=0.017$), but also
234 dose reduction due to AE ($P=0.003$) and discontinuations because of AEs ($P=0.035$). Additionally, Cox
235 proportional hazards modeling suggested a higher risk of ILD with higher exposure of intact DS-
236 8201a (any grade, $P<0.001$; grade ≥ 2 , $P=0.007$). Based on the predicted benefit / risk profile, 5.4mg/kg
237 DS-8201a was chosen as the recommended dose for further development of DS-8201a in HER2-
238 positive breast cancer.

239 Thirty-four patients with HER2-low breast cancer, were enrolled at data cutoff of October 12 2018,
240 with a median age of 56 (range 33-75) and a median number of prior cancer regimens of 8 (2-18) [33].
241 Most of the HER2-low patients had hormone-receptor positive disease (87%) and 34% of them were
242 pretreated with a CDK4/6 inhibitor. The confirmed ORR was 44%, the DCR was 79% and the median
243 time to response was 2.8 months (range 1.6 – 3.0 months). Median PFS was 7.6 months (95% CI 4.9-
244 13.7) and duration of response (DOR) was 9.4 months (95% CI 1.5-23.6). In a subgroup analysis based
245 on IHC expression of HER2, ORR (54% vs 33%) and median PFS (13.6 months vs 5.7 months) were
246 superior in IHC 2+ tumors ($N=24$) compared to IHC 1+ tumors ($N=27$), respectively.

247 **Ongoing trials without published results.** Currently 7 registered trials investigating DS-8201a are
248 active. In a phase 1b trial with a dose escalation and an expansion cohort, the combination of DS-
249 8201a with the PD-1 checkpoint-inhibitor nivolumab in patients with breast and urothelial
250 carcinomas is studied (clinicaltrials.gov identifier: NCT03523572). One phase 1 trial assess the safety
251 and pharmacokinetics in HER2-positive advanced gastric cancers, gastroesophageal junction
252 adenocarcinomas or breast cancers (clinicaltrials.gov identifier: NCT03368196). A third phase 1 trial
253 investigates the effect on QT intervals and pharmacokinetics of different DS-8201a doses in patients
254 with HER2-positive breast cancer (clinicaltrials.gov identifier: NCT03366428). In the phase 2 trial
255 DESTINY-Breast01, HER2-positive breast cancer patients resistant, refractory or intolerant to T-DM1
256 are randomized into different DS-8201a dose level groups, to assess pharmacokinetics and
257 recommended dose, followed by an expansion cohort (clinicaltrials.gov identifier: NCT03248492).

258 The randomized, open-label phase III trial DESTINY-Breast02 investigates DS-8201a compared to
259 treatment of physicians's choice (trastuzumab plus capecitabine or lapatinib plus capecitabine) in
260 patients with HER2-positive advanced breast cancer (ABC) pretreated with prior standard of care
261 HER2 therapies including T-DM1 (clinicaltrials.gov identifier: NCT03523585). The two ADCs DS-
262 8201a and T-DM1 are compared in the randomized, open-label phase III trial DESTINY-Breast03 in

263 pretreated patients with HER2-positive ABC (clinicaltrials.gov identifier: NCT03529110). The third
264 ongoing randomized, open-label phase 3 trial (DESTINY-Breast04), studies DS-8201a compared to
265 treatment of physicians's choice (capecitabine, eribulin, gemcitabine, paclitaxel or nab-paclitaxel) in
266 patients with HER2-low (IHC 1+ or IHC 2+ in situ hybridization [ISH]-) ABC (clinicaltrials.gov
267 identifier: NCT03734029).

268

269 7. MEDI4276

270 **Drug Structure.** MEDI4276 is a novel ADC composed of a HER2-bispecific antibody targeting two
271 different epitopes on HER2, site-specific conjugated via a maleimidocaproyl linker to the potent
272 tubulysin-based microtubule inhibitor AZ13599185 [34,35]. The small-molecule toxin AZ13599185 is
273 a microtubule polymerization inhibitor during mitosis inducing cell death. The bispecific antibody
274 contains four antigen-binding units, two on each arm that are capable of interacting with two
275 different epitopes on HER2. Antibody interaction with the unique HER2 epitope, different to the
276 trastuzumab and pertuzumab binding epitope, can completely interfere with HER2-HER3 receptor
277 dimerization induced by heregulin-1 β . Therefore, this bispecific antibody blocks both ligand-
278 independent and ligand-dependent receptor activation. The DAR of MEDI4276 is 4.

279 **Preclinical data.** *In vitro*, MEDI4276 was at least 10-fold more potent in the HER2 overexpressing cell
280 line SKBR-3, than T-DM1 [34]. Additionally, MEDI4276 demonstrated efficacy in the T-DM1 resistant
281 HER2 positive JIMT-1 cell line. In HER2-low cell lines (MCF7-GTU, and ZR-75-1), MEDI4276 revealed
282 potent cell killing whereas T-DM1 was inactive. In a patient derived HER2-positive breast cancer
283 xenograft model, weekly intravenous administration of MEDI4276 over four weeks induced a
284 complete remission in all treated animals which retained tumor free over 120 days after treatment. In
285 contrast, T-DM1 only induced tumor stasis and a regrowth was observed soon after T-DM1 treatment
286 was stopped. In a several patient derived HER2-low xenograft models, MEDI4276 induced tumor
287 regression regardless of the hormone receptor status.

288 **Clinical data.** In a phase 1/2 dose escalation and dose expansion trial MEDI4276 was investigated in
289 patients with advanced pretreated HER2 expressing (IHC 2+) breast or gastric cancer
290 (clinicaltrials.gov identifier NCT02564900) [36]. As of November 1 2017, 43 patients were enrolled
291 and treated in several dose cohorts (0.05, 0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.75, or 0.9 mg/kg every 3 weeks),
292 following a 3+3 design. MTD was exceeded at 0.9 mg/kg. Drug-related AEs of any grade and grade \geq
293 3 were reported in 88% and 12% of patients, respectively. The most common AEs were nausea (all
294 grade: 58%), fatigue (all grade: 42%), elevated AST (all grade: 37%, grade \geq 3: 19%), vomiting (all
295 grade: 37%), and elevated ALT (all grade: 35%, grade \geq 3: 12%). Drug-related peripheral neuropathy
296 grade 3 was observed in 1 patient (2%) at 0.6 mg/kg and in 2 patients (5%) at 0.75 mg/kg. In evaluable
297 patients 1 CR (0.5 mg/kg; breast cancer), 1 PR (0.6 mg/kg; breast cancer), and 12 SD (28%) were
298 reported. A non-linear pharmacokinetic with rapid clearance and negligible deconjugation of
299 MEDI4276 was observed.

300 The preclinical data and early clinical data of MEDI4276 supports further clinical development of this
301 drug in HER2-positive and HER2-low breast cancer patients.

302

303 8. RC48 (RC48-ACD, hertuzumab-vc-MMAE)

304 **Drug Structure.** RC48 is a novel humanized anti-HER2 antibody hertuzumab conjugated with
305 monomethyl auristatin E (MMAE) via a cleavable linker [37]. MMAE acts by inhibiting the tubulin
306 polymerization in dividing cells resulting in an in G2/M phase arrest and apoptosis [23]. Hertuzumab

307 had a higher affinity to HER2 than trastuzumab in an ELISA-based binding assay. The monoclonal
308 anti-HER2 antibody binds specifically to HER2, but not to other members of the human epidermal
309 growth factor receptor family (EGFR, HER3, or HER4). After binding of hertuzumab to HER2-
310 expressing tumor cells on the cell surface, fluorescence-labeled hertuzumab is internalized through
311 endocytosis, later detectable in lysosomes. MMAE is linked to hertuzumab using a protease-sensitive
312 valine-citrulline dipeptide sequence, which was designed for optimal stability in human plasma and
313 efficient cleavage by human cathepsin B. The DAR is approximately 4. After binding of RC48 to HER2
314 on the cell surface, the R48-HER2 complex is internalized through endocytosis. Following
315 internalization, lysosomal proteases cleave both, the monoclonal antibody and the linker, and MMAE
316 is released.

317 **Preclinical data.** In vivo efficacy of RC48 was investigated in trastuzumab and lapatinib sensitive
318 and resistant breast cancer xenograft models in female nude BALB/cA mice subcutaneously
319 inoculated with breast cancer cells [37]. For the trastuzumab and lapatinib sensitive model, BT-474
320 human breast cancer cells, which express high levels of HER2, were implanted. Antitumor activity of
321 RC48 (0.5, 1.5, and 5.0 mg/kg) was dose-dependent. RC48 activity at dose levels ≥ 0.5 mg/kg was
322 significantly stronger compared to trastuzumab (10 mg/kg) and lapatinib (200 mg/kg). In the resistant
323 breast cancer model, nude mice were inoculated with BT-474/T721 (trastuzumab-resistant) and BT-
324 474/L1.9 (trastuzumab- and lapatinib-resistant) cells, respectively. Both RC48 (5.0 mg/kg) and T-DM1
325 (5.0 mg/kg) were significantly more efficacious in the BT-474/T721 xenograft model compared to
326 trastuzumab. In the trastuzumab and lapatinib resistant BT474/L1.9 xenograft model, RC48 (5.0
327 mg/kg) was significantly more effective than trastuzumab, lapatinib and T-DM1.

328 **Clinical data.** RC48 is investigated in a dose escalation open-label, single-center phase 1 trial in
329 HER2-positive breast cancer patients (clinicaltrials.gov identifier: NCT02881138) [38]. As of January
330 29 2018, 23 patients were treated in 5 dose escalation cohorts (dose levels 0.5, 1.0, 1.5, 2.0, 2.5 mg/kg)
331 once every two weeks (Q2W) following a 3+3 design. Median age was 57 years (range 32-65), median
332 prior treatment lines in the metastatic setting was 3 (range 1-6), and 70% (16/23) of patients were
333 pretreated with trastuzumab. MTD was not reached at doses up to 2.0 mg/kg Q2W. The most
334 common treatment-related adverse events (AEs) were leucopenia (all grades: 48%, grade ≥ 3 : 4%),
335 AST elevation (all grades: 48%, grade ≥ 3 : 4%) and neutropenia (all grades: 43%, grade ≥ 3 : 13%). In
336 patients treated at doses ≥ 1.5 mg/kg (14 evaluable patients for response), ORR was 57% (8/14) and
337 DCR was 86% (12/14). Because MTD was not determined at dose levels up to 2.0 mg/kg twice-weekly,
338 a 2.5 mg/kg Q2W dose escalation cohort is ongoing.

339 A second dose escalation phase 1 trial investigates RC48 in patients with HER2-overexpressing
340 advanced solid cancers (clinicaltrials.gov identifier: NCT02881190) [39]. As of January 29 2018, 36
341 patients, including 1 breast cancer patient, were enrolled in dose escalation (0.1 - 2.5 mg/kg Q2W and
342 2.0 mg/kg Q3W) and dose expansion cohorts, respectively. The most common treatment-related AEs
343 were in line with the previously mentioned trial: AST elevation (all grades: 50%, grade ≥ 3 : 3%), ALT
344 elevation (all grades: 43%, grade ≥ 3 : 3%), leucopenia (all grades: 33%, grade ≥ 3 : 7%), neutropenia (all
345 grades: 33%, grade ≥ 3 : 10%) and numbness (all grades: 23%, grade ≥ 3 : 0%). No AEs grade ≥ 4 were
346 observed. Pharmacokinetic analyses demonstrated a dose-dependent exposure with a 1-1.5 days half-
347 life [40]. A further expansion cohort investigating a dose of a 2.5 mg/kg is planned.

348 An open-label, multicenter phase 1b/2 trial investigates RC42 in patients with pretreated metastatic
349 HER2-positive breast cancer (clinicaltrials.gov identifier: NCT03052634) [40]. As of January 2018, 30
350 patients (6 IHC 2+/ISH+; 24 IHC 3+) were enrolled in 1.5 and 2.0 mg/kg cohorts in the phase 1b part
351 of the trial. Median age was 53 years (range 26-62), 19 patients (63%) were pretreated with HER2-
352 targeting drugs and 16 patients (53%) were pretreated with ≥ 3 prior chemotherapy regimens in the
353 metastatic setting. ORR was 37% (11 PR) and DCR was 97% (29/30) with a clinical benefit rate (CBR;
354 CR + PR + SD ≥ 6 months) of 47% (14/30). In the 1.5 mg/kg and 2.0 mg/kg cohorts the ORR was 27%
355 and 47%, respectively. In trastuzumab-naive and trastuzumab-pretreated patients ORR were 57%

356 and 33%, respectively. Most common treatment-related AEs were in line with the two previously
357 mentioned RC42 phase 1 trials: AST elevation (all grades: 50%, grade \geq 3: 3%), ALT elevation (all
358 grades: 43%, grade \geq 3: 3%), leucopenia (all grades: 33%, grade \geq 3: 7%), neutropenia (all grades: 33%,
359 grade \geq 3: 10%), numbness (all grades: 23%, grade \geq 3: 0%). Thrombocytopenia (all \leq grade 2) was
360 observed in 10% of patients. No grade \geq 4 AEs were observed. Enrollment in the 2.5 mg/kg expansion
361 cohort is underway. In the planned phase 2 part of the trial, patients will be randomized to RC48 at
362 the dose level selected in phase 1b or to lapatinib plus capecitabine.

363 **Ongoing trials without published results.** A randomized, multicenter, 2-arm, open-label phase II
364 trial comparing RC48 (2.0 mg/kg Q2W) with capecitabine plus lapatinib in trastuzumab pretreated
365 patients with advanced HER2-positive breast cancer is currently recruiting in Chinese trial centers
366 (clinicaltrials.gov identifier: NCT02881138).

367 The novel ADC RC42 demonstrated a favorable toxicity profile in 3 phase I trials. Response rates of
368 37-57% in partly heavily pretreated patients with HER2-positive breast cancer patients is promising
369 and further clinical development of this drug is warranted.

370

371 9. SYD985 ([vic-]trastuzumab duocarmazine)

372 **Drug structure.** SYD985 is composed of the monoclonal HER2 directed antibody trastuzumab linked
373 via a cleavable valine-citrulline peptide to the synthetic duocarmycin analogon seco-DUocarmycin-
374 hydroxyBenzamide-Azaindole (vc-seco-DUBA) [41,42]. Duocarmycins are DNA-alkylating agents
375 composed of a DNA-alkylating and a DNA-binding moiety binding into the minor groove of the
376 DNA causing irreversible alkylation of DNA [42]. These cytotoxic drugs induce cell death in both
377 dividing and nondividing cells by disrupting the nucleic acid architecture. The average DAR is 2.8.

378 **Preclinical data.** In a trastuzumab sensitive BT474 mouse xenograft model, antitumor activity of
379 SYD985 was dose depended [42]. Antitumor activity of 1 mg/kg SYD985 was equal to 5 mg/kg
380 trastuzumab and SYD985 dosed once at 5 mg/kg significantly reduced tumor growth compared to
381 trastuzumab at the same dose level. In HER2-positive (IHC 3+) breast cancer patient derived
382 xenograft models named MAXF1322 and MAXF1162, SYD985 dose dependently reduced tumor
383 growth, whereas high dose trastuzumab did not reveal any antitumor activity. In a HER2-positive
384 (IHC 3+) breast cancer cell line (SK-BR-3) and trastuzumab-resistant breast cancer cell line (UACC-
385 893), SYD985 and T-DM1 demonstrated similar potencies [43]. In two HER2-low (HER 1+) cell lines
386 (MDA-MB-175-VII and ZR-75-1), SYD985 retained its activity, whereas T-DM1 was less potent.
387 Neither SYD985 nor T-DM1 were able to kill HER2-negative cells (SW-620 and NCI-H520). These
388 findings were confirmed *in vivo* where SYD985 was active in HER2-low breast cancer xenograft
389 models in contrast to T-DM1. In coculture experiments of HER expressing cells (SK-BR-3 and MDA-
390 MB-175-VII) with HER2 negative (HER2 0) NCI-H520 cells, a bystander killing was observed in
391 presence of SYD985, but not of T-DM1.

392 **Clinical data.** SYD985 was investigated in a two-part phase 1 trial (clinicaltrials.gov identifier:
393 NCT02512237). In the dose-escalation part of the study, patients with solid tumors and any HER2
394 status (n=39), including 26 patients with breast cancer, were enrolled and treated with SYD985 doses
395 varying from 0.3 mg/kg to 2.4 mg/kg every three weeks. The RP2D was defined as 1.2 mg/kg Q3W.
396 Patients with HER2 expressing breast, gastric, urothelial or endometrial cancers were subsequently
397 enrolled in expansion cohorts treated with the RP2D. Breast cancer patients (n=26) enrolled in the
398 dose-escalation part were heavily pretreated with a median of 7 systemic therapies [44]. All HER2-
399 positive patients were pretreated with T-DM1. As of May 16 2016, tumor evaluation data were
400 available for 19 of 26 enrolled breast cancer patients. In evaluable HER2-positive patients (n=14), ORR
401 was 36% (5/14) and DCR was 93% (13/14). In evaluable HER2-low patients (n=5) an ORR of 60% (3/5)

402 and a DCR of 80% (4/5) was observed. In evaluable patients treated with doses ≥ 1.2 mg/kg ORR were
403 42% and 75% for the HER2-positive and HER2-low patients, respectively. One fatal pneumonitis
404 occurred at 2.4 mg/kg of SYD985. Up to doses of 1.8 mg/kg every 3 weeks, SYD985 was well tolerated.
405 The most frequently reported drug-related AEs were conjunctivitis, stomatitis, fatigue, and decreased
406 appetite and the majority of these AEs were of mild or moderate intensity.

407 Ninety-nine breast cancer patients were enrolled in dose expansion cohorts: 50 patients with HER2-
408 positive MBC, 32 patients with HER2-low hormone-receptor positive disease and 17 patients with
409 HER2-low triple negative MBC [45]. The median number of prior cancer regimens was 6 (range 1-21)
410 and 80% of the HER2-positive patients were pretreated with T-DM1. In HER-positive patients ORR
411 was 33% (16/48 patients with measurable disease) and PFS was 9.4 months (95% CI 4.5-12.4). In T-
412 DM1 pretreated HER2 positive patients, an ORR of 29% (11/38) and a PFS of 8.3 months (95% CI 4.1-
413 15.0) was observed. ORR was 27% (8/30) and 40% (6/15), and PFS was 4.1 (95% CI 2.4-5.4) and 4.4
414 (95% CI 1.0-7.1) in patients with HER2-low hormone-receptor positive and HER2-low triple negative
415 disease, respectively. The most common drug related AEs in patients of all expansion cohorts (n=146)
416 were fatigue (all grades: 32%, grade ≥ 3 : 3%), dry eyes (all grades: 29%, grade ≥ 3 : 1%), conjunctivitis
417 (all grades: 25%, grade ≥ 3 : 3%) and nausea (all grades: 20%, grade ≥ 3 : 0%). The majority of AEs were
418 grade 1 or 2 with 6% of grade 3 AEs. No \geq grade 4 AEs were observed. Twenty-eight (19%) of patients
419 discontinued treatment due to AEs, most commonly due to ocular toxicity. Alopecia was reported in
420 18% of patients (grade 1: 15%, grade 2: 3%).

421 **Ongoing trials without published results.** SYD985 currently investigated in a multi-center, open-
422 label, randomized phase 3 trial comparing SYD985 with physician's choice in patients with HER2-
423 positive advanced or metastatic breast cancer pretreated with T-DM1 (clinicaltrials.gov identifier:
424 NCT03262935).

425 SYD985 was well tolerated and ocular toxicity was commonly reported in a large phase 1 trial. The
426 efficacy data of a phase 1 expansion cohort in T-DM1 pretreated patients with HER2 positive breast
427 cancer are promising. The results of an ongoing phase 3 trial in this patient population is awaited
428 within the next two years.

429

430 **10 XMT-1522 (TAK-522)**

431 **Drug structure.** XMT-1522 is an ADC composed of a novel IgG1 anti-HER2 monoclonal antibody
432 (HT-19) conjugated with the Dolaflexin® platform to auristatin-based drug payload molecules
433 (Auristatin F-hydroxypropylamide, AF-HPA) [46,47]. The Dolaflexin® platform is a biodegradable
434 polymer-based conjugation platform that enables a high average XMT-1522 DAR of 12 (range 10-15)
435 without aggregation or detrimental impact on pharmacokinetics [48]. Auristatin analogs act by
436 inhibiting the tubulin polymerization in dividing cells resulting in an in G2/M phase arrest and
437 apoptosis [23]. The HT-19 antibody is non-competitive with trastuzumab or pertuzumab for HER2
438 binding.

439 **Preclinical data.** Across a panel of 25 tumor cell lines with different HER2 expression levels, XMT-
440 1522 was approximately hundred times more potent than T-DM1 [46]. In a BT-474 HER2-positive
441 breast cancer xenograft model, a single dose of 5 mg/kg HT-19 antibody was inactive, while a single
442 dose of 2 mg/kg or 5 mg/kg XMT-1522 induced durable complete tumor regression, indicating that
443 the primary mechanism of XMT-1522 is cytotoxic payload delivery, not HER2 signaling inhibition.
444 In the same model, T-DM1 at a single dose of 5 mg/kg was inactive. In a patient-derived HER2-
445 positive xenograft model, XMT-1522 induced a durable complete tumor regression after a single 1
446 mg/kg dose, while a 10 mg/kg dose of T-DM1 achieved a tumor growth delay without regression

447 only. In a patient-derived HER2-low xenograft model, XMT-1522 at a single 3 mg/kg dose achieved
448 a partial tumor regression, whereas T-DM1 was inactive.

449 *In vitro*, a combination of XMT-1522 with trastuzumab did not block the XMT-1522 HER2 binding
450 ability or the ADC internalization. In a HER2 positive xenograft model, a combination of
451 trastuzumab, pertuzumab and XMT-1522 was synergistic. Despite the high potency of XMT-1522 in
452 HER2-low tumor models, no XMT-1522-related toxicity was observed in HER2-expressing tissues
453 including heart and lung [49].

454 In multiple cell lines, an immunogenic cell death, as measured by cell surface expression of
455 calreticulin, was induced a few hours after treatment with free AF-HPA and XMT-1522, respectively
456 [50]. In a HER2-low breast cancer (4T1) xenograft model, XMT-1522 but not T-DM1 significantly
457 inhibited tumor growth. A combination of XMT-1522 with an anti-PD1 monoclonal antibody
458 synergistically enhanced the anti-tumor efficacy, with complete responses in some mice. The
459 frequency of complete remissions was further enhanced when the two drugs were given sequentially
460 (XMT-1522 followed by the checkpoint inhibitor).

461 **Clinical data.** XMT-1522 is currently investigated in the first-in-human phase 1b dose escalation and
462 expansion trial in patients with advanced HER2-expressing (IHC $\geq 1+$) breast cancer, gastric cancer
463 and non-small cell lung cancer progressing on standard therapy (clinicaltrials.gov identifier:
464 NCT02952729). XMT-1522 is administered intravenously every 3 weeks. Dose escalation uses a 3+3
465 design and a 3-week dose limiting toxicity (DLT) evaluation period. As of February 1 2018, 19 patients
466 have completed the DLT evaluation period across 6 dose levels (2 to 21.3 mg/m² every 3 weeks). Since
467 no DLT, no serious adverse event (SAE) and no treatment-related AE \geq grade 3 have been observed,
468 dose escalation was continued [51]. The most common treatment-related AE were elevated liver
469 enzymes, fatigue, nausea, vomiting, headache, and anorexia. ORR was 17% (1/5 evaluable patients)
470 and DCR was 83% (5/6) in patients dosed at 16 or 21.3 mg/m². The partial remission was observed at
471 the first assessment in a patient with HER2-positive breast cancer previously treated with T-DM1. In
472 patients treated at doses less than 16 mg/m², DCR was 25% (3/12) with no observed responses.
473 Systemic exposure of total AF-HPA payload was approximately dose-proportional. Plasma
474 concentrations of free AF-HPA and its active metabolites were low.

475 XMT-1522 has interesting biochemical features with a higher drug-antibody ratio and a novel HER2
476 antibody. Preclinical data and first clinical data are promising and the final results of the first-in-
477 human phase 1 trial are awaited within the next year.

478

479 11. Discussion

480 Antibody-drug conjugates are a promising class of anti-cancer drugs combining the selectivity of
481 monoclonal antibodies and the cell killing potential of cytotoxic agents [15,52]. Targeted cytotoxic
482 drug delivery into tumor tissue increases the therapeutic window of these agents considerably. For
483 example, clinical development of unconjugated DM1 was stopped early due to unfavorable toxicity
484 despite promising clinical activity [52]. In contrast T-DM1, consisting of DM1 attached via a non-
485 cleavable linker to trastuzumab, has a favorable toxicity profile and a clinical meaningful antitumor
486 effectivity [17].

487 The published preclinical and clinical data of the reviewed investigational HER2 directed ADCs
488 A166, ALT-P7, ARX788, DHES0815A, DS-8201a, RC48, SYD985, MEDI4276 and XMT-1522 are
489 promising. In preclinical models, most of these drugs were more effective than T-DM1, which raises
490 high expectations for these novel drugs. The investigation of SYD985 and DS-8201a in T-DM1-
491 refractory HER2-positive patients in currently enrolling randomized phase 3 trials is a very favorable

492 development. Notably, the addition of new treatment option, instead of replacing an established
493 option by an equally or more effective drug, has been shown to have a greater impact on survival of
494 metastatic breast cancer patients [53,54].

495 Which out of the plethora of new ADCs will find its way into the clinic, remains speculative.
496 Interestingly, the toxicity profile of the different compounds is quite different, which could influence
497 patient and physician's choice in case of comparable efficacy. All ADCs show some hematologic and
498 hepatic toxicity, however DS-8201a, harboring a topoisomerase I inhibitor, showed additional
499 gastrointestinal toxicity, while SYD985, revealed an unfamiliar ocular toxicity. How these toxicities
500 will influence treatment intensity and adherence, future phase II and phase III trials will uncover.

501 Preclinical and early clinical efficacy data of DS-8201a, SYD985, MEDI4276 and XMT-1522 in HER2-
502 low breast cancers is of special interest. About 50% of breast cancers can be categorized as HER2-low
503 and the availability of a targeted treatment option for this patient population would be of a great
504 interest. This is especially true for patients with triple-negative breast cancer, the breast cancer
505 subtype with the worst prognosis for whom still no targeted treatment options are available.

506 It can be expected that the evidence of clinical efficacy of these promising novel HER2 directed ADCs
507 will increasingly corroborate.

508

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526 **Abbreviations**

ABC	advanced breast cancer
ADC	antibody-drug-conjugates
AE	adverse event
AF-HPA	auristatin f-hydroxypropylamide,
ALT	alanine transaminase
AST	aspartate transaminase
CBR	clinical benefit rate
CI	confidence interval
CR	complete response
DAR	drug antibody ratio
DCR	disease control rate
DFS	disease-free survival

DOR	duration of response
EGF	epidermal growth factor
EGFR	epidermal growth factor receptor
HER2	human epidermal growth factor receptor 2
HER3	human epidermal growth factor receptor 3
HER4	human epidermal growth factor receptor 4
ISH	<i>in situ</i> hybridization
IHC	immunohistochemistry
ILD	interstitial lung disease
MBC	metastatic breast cancer
MDT	maximum tolerated dose
MMAE	monomethyl auristatin E
MMAF	monomethyl auristatin F
ORR	overall response rate
PFS	progression-free survival
PBD-MA	pyrrolo[2,1- c][1,4]benzodiazepine monoamide
PR	partial response
Q2W	every 2 weeks
Q3W	every 3 weeks
ROR2	receptor tyrosine kinase-like orphan receptor 2
RP2D	recommended phase 2 dose
SAE	serious adverse event
SD	stable disease

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