Cysteine in cell culture media induces acidic IgG1 species by disrupting the disulfide bond network

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Funding information
Boehringer Ingelheim

Abstract
A high degree of charge heterogeneity is an unfavorable phenomenon commonly observed for therapeutic monoclonal antibodies (mAbs). Removal of these impurities during manufacturing often comes at the cost of impaired step yields. A wide spectrum of posttranslational and chemical modifications is known to modify mAb charge. However, a deeper understanding of underlying mechanisms triggering charged species would be beneficial for the control of mAb charge variants during bioprocessing. In this study, a comprehensive analytical investigation was carried out to define the root causes and mechanisms inducing acidic variants of an immunoglobulin G1-derived mAb. Characterization of differently charged species by liquid chromatography–mass spectrometry revealed the reduction of disulfide bonds in acidic variants, which is followed by cysteinylation and glutathionylation of cysteines. Importantly, biophysical stability and integrity of the mAb are not affected. By in vitro incubation of the mAb with the reducing agent cysteine, disulfide bond degradation was directly linked to an increase of numerous acidic species. Modifying the concentrations of cysteine during the fermentation of various mAbs illustrated that redox potential is a critical aspect to consider during bioprocess development with respect to charge variant control.

KEYWORDS
acidic charge variants, bioprocess optimization, cysteine, product quality, redox-sensitive modifications

1 | INTRODUCTION

With more than 300 molecules currently in development for diverse medical applications, monoclonal antibodies (mAbs) have emerged as the leading class of biopharmaceuticals (Shukla et al., 2017). Product heterogeneity of therapeutic mAbs is commonly observed, as they are exposed to numerous factors during bioprocessing, such as media components, fermentation conditions and downstream processing (Gandhi et al., 2012; Hossler et al., 2015; Kishishita et al., 2015; Xie et al., 2016). These conditions result in product-related impurities, which must be closely controlled and monitored during the manufacturing process, as mandated by regulatory authorities (ICH Q 6 B).

Charge heterogeneity in mAbs is considered a critical quality attribute, as it can alter efficacy and pharmacokinetics (Boswell et al.,...
Unpaired cysteines can originate from insufficient formation of disulfide bridges during protein folding and assembly, as well as the degradation of intact bonds. A higher degree of free thiols is reported for cysteines involved in intrachain as compared to interchain disulfide bonds (Chumsae et al., 2009; Xiang et al., 2009). Cysteines involved in intrachain bonds are protected from degradation as they are buried between the β-sheets of the IgG fold, indicating that these unpaired cysteines must originate from incomplete formation during the production process (Amzel & Poljak, 1979; Kikuchi et al., 1986). In particular, the redox environment during fermentation may be critical for a correct disulfide bond network (Chaderjian et al., 2005; Handlogten et al., 2017; Kshirsagar et al., 2012). This is supported by the observation that the addition of copper sulfate to cell culture facilitates the formation of an intact intrachain disulfide bridge by acting as an oxidative agent (Chaderjian et al., 2005). Accordingly, interchain disulfide bonds are reduced by the thioredoxin and glutathione pathway during fermentation (Handlogten et al., 2017).

The importance of controlling disulfide bridge conversions and resulting charge variants during bioprocessing is highlighted by reports where a decrease in potency has been linked to free thiols, in particular in the variable domains (Harris, 2005; Ouellette et al., 2010). Destabilizing effects of incomplete disulfides have been observed for some cases, but they do not introduce substantial changes to the IgG structure (Kikuchi et al., 1986; McAuley et al., 2008). Although there is general consensus on chemical and structural modifications contributing to charge variants, there is a lack of understanding of the underlying mechanisms responsible for acidic variants (Chung et al., 2018). Previously published reports comparing CEX fractions by bioanalytical techniques have shown that the origin of mAb charge variants is heterogeneous and that combinations of protein modifications can complicate their characterization (Griaud et al., 2017; Miao et al., 2017; Ponniah et al., 2015). This study aimed to fully characterize the root cause of charged species of an IgG1 mAb. By characterizing individual charged mAb species, the

![Schematic representation of an immunoglobulin G1 (IgG1) half molecule. Domains are labeled in bold and cysteines are shown in italic. The antigen-binding fragment (Fab) consists of the variable (V) and constant (C) domains of the light chain, and the variable (V) and constant (C) domains of the heavy chain. The Fab fragment is connected via the hinge region to the crystallizable fragment (Fc), consisting of the constant domains C and C of the heavy chain. A total of 16 disulfide bonds stabilize the native IgG1 structure, as represented by dashed lines connecting the cysteines involved](image-url)
generation of acidic mAb1 species was associated to ruptured and modified disulfide bonds. These modifications are induced by cysteine, as demonstrated by in vitro experiments. In cultivation experiments, it was shown that the media component cysteine triggers changes in mAb charge profile and alters the media redox potential. The cysteine concentration in cell culture media may thus be crucial for the control of mAb charge during bioprocessing.

2 MATERIALS AND METHODS

2.1 Materials

All mAbs investigated in this study are IgG1 with pI of 8.5–8.9. mAb1, mAb2, and mAb3 are IgG1 and mAb4 is an IgG1-derived antibody, all produced in Chinese hamster ovary (CHO) cells. All chemicals and reagents used in this study were high performance liquid chromatography (HPLC) grade purchased from Sigma-Aldrich Chemie GmbH.

2.2 mAb1 purification by protein A affinity chromatography

mAb1 was purified from cell culture free fluid using an in-house-packed MabSelect SuRe column, operated on an ÄKTA avant 25 using UNICORN 7.0 (GE Healthcare GmbH). The protein was eluted in 50 mM acetate pH 3.5. The eluted product pool was neutralized to pH 5.6 and stored at −70 °C until further use.

2.3 Treatment of mAb1 samples with cysteine, S-sulfocysteine, cystine, and glutathione

Intact and fractionated mAb1 were incubated with cysteine (n = 4), S-sulfocysteine (n = 3), and cystine (n = 3) at a protein concentration of 2 g/L in 10 mM acetate at pH 5.6. Glutathione incubations of intact mAb1 with 2.5 mM GSH and 2.5 mM GSSG (n = 2) were carried out at a protein concentration of 2 g/L in 50 mM Tris at pH 8.2. Cysteine, S-sulfocysteine, GSH, and GSSG stock solutions were prepared in Milli-Q® (Merck-Millipore) at 100X and added to the mAb to reach the respective final concentrations. Before further analysis, the samples were rebuffered into 10 mM sodium phosphate pH 6.8 using 7 kDa molecular weight cut-off Zeba™ Spin Desalting Columns (Thermo Fisher Scientific Inc.) to remove the reagents.

2.4 Preparation of mAb1 charge variants

Preparative separation of the mAb charge variants was carried out on a 9 × 250 mm ProPac™ WCX-10 (Thermo Fisher Scientific Inc.) using an ÄKTA avant 25 and UNICORN 7.0 (GE Healthcare GmbH). mAb1 fractions were eluted using a linear NaCl gradient and pooled to obtain the charge variants as shown in Figure S1. As a control, all fractions of one run were pooled to obtain a sample of all charge variants exposed to the same conditions as the fractions. The charge variants and control sample were concentrated to around 2–5 g/L and rebuffered to 10 mM sodium phosphate pH 6.8 using 10 kDa molecular weight cut-off Amicon® Ultra-15 centrifugal filter units (Sigma-Aldrich Chemie GmbH). In total, the fractionation experiment was carried out twice, and all resulting samples were checked by weak cation exchange chromatography (WCX) and capillary electrophoresis (CE). Biophysical properties (circular dichroism [CD], nano differential scanning fluorimetry [nano-DSF], size exclusion chromatography [SEC]) were determined from one run in duplicates.

2.5 Capillary electrophoresis (CE)

Protein size variants were quantified via microfluidic CE using the LabChip GXII instrument and LabChip GX 5.3 software (PerkinElmer). Samples were prepared according to the manufacturer’s specifications of the Protein Express Assay Reagent Kit, and separated on Protein Express Assay LabChip using the HT Protein Express 200 run (PerkinElmer).

2.6 Analytical separation and quantification of charge variants

Analytical separation and quantification of mAb1, mAb2, and mAb3 charge variants was achieved by analytical CEX, operated on Alliance HPLC systems (Waters). mAb1 and mAb3 were separated on a 4 × 250 mm ProPac WCX-10 column and mAb2 was separated on a 4 × 250 mm MabPac™ SCX-10 column (Thermo Fisher Scientific Inc.). System control and data analysis was performed using EMPOWER 3 (Waters), mAb4 charge variant analysis was performed by capillary zone electrophoresis using a PA 800 Plus (Beckman Coulter).

2.7 Analytical size-exclusion chromatography (SEC)

Analytical separation and quantification of mAb1 high molecular weight (HMW), intact (main), and low molecular weight species (LMW) was achieved on a 4.6 × 30 mm TSKgel UP-SW3000 (Tosoh Bioscience LLC) using an Acquity UPLC system (Waters). System control and data analysis was performed using EMPOWER 3 (Waters).

2.8 Thermal unfolding by nano-differential scanning fluorimetry (nano-DSF)

Protein stability was assessed via thermal unfolding detected by nano-DSF on the Prometheus NT.48 (NanoTemper Technologies GmbH). Protein samples were measured using a temperature ramp from 45°C to 85°C. System control and data analysis were performed using PR.ThermControl (NanoTemper Technologies GmbH).
2.9 | Far UV CD spectroscopy

Far UV CD measurements were acquired on a JASCO spectrometer at protein concentrations ranging from 1.5 to 2.0 g/L. Spectra were recorded in millidegrees (m°).

2.10 | Combined endopeptidase Lys-C/tryptic digest with LC-MS/MS analysis for quantification of free thiols, cysteinylated, and glutathionylated thiols

Samples were subjected to a combined Lys-C/tryptic digest as described previously (Seibel et al., 2017). For analysis of free thiols, half of the Lys-C/tryptic digest was reduced by addition of 4.4 µl 0.64 M Tris-(2-carboxyethyl)-phosphine (Roth) and incubation for 30 min at 37°C. Subsequently, 8 µl 20% formic acid to stop the digestion and analyzed by liquid chromatography–mass spectrometry (LC-MS/MS).

Sample measurement was conducted as described previously (Seibel et al., 2017). For analysis of disulfide connectivity, reduced and nonreduced data sets from the same sample were analyzed using PepFinder™ (Thermo Fisher Scientific Inc.) and manually reviewed (n = 1). Extracted ion chromatograms (EICs) of each dipeptide were generated using the two or three most intense charge states. The peak areas of all EIC peaks containing disulfide linked peptides were set to 100% and the relative amount of each disulfide linked total peak area of all EIC peaks containing disulfide linked peptides was calculated serving for inter-sample comparison. "Free" thiols were relatively quantified using the peak areas of the EICs of the NEM-modified ("free thiol") peptide and the reduced peptide as 100% value. Cysteinylated and glutathionylated peptides were quantified using the nonreduced peptide map data and the EIC peak areas of NEM-alkylated ("free thiol"), the cysteinylated, the glutathionylated peptide and dipeptide as 100% value. For relative quantification of "free" cysteines, cysteine-containing tryptic peptides were selected that showed good chromatographic and mass spectrometric properties for both, the alkylated and the non-alkylated cysteine-containing tryptic peptide.

2.11 | LC-MS analysis of deglycosylated intact, reduced and IdeS digested samples

For determination of the intact mass (n = 1), the antibody was deglycosylated with PNGaseF (R&D Systems) at 37°C over night according to manufacturer’s instructions. For measurement of the reduced antibody samples, deglycosylated samples were treated with 8 M guanidine hydrochloride and Tris-(2-carboxyethyl)phosphine hydrochloride for 30 min at 37°C. For cleavage in the hinge region, antibody samples were incubated with IdeS enzyme (Fabricator, Genovis) for 2 h at 37°C according to manufacturer’s instructions. Samples were analyzed by LC-MS using a linear organic solvent gradient on C4 column (Acquity BEH300 C4, 1 mm, 50 mm, 1.7 µm; 150 µl/min, 75°C, 1.6 µg on column) and a Quadrupole time-of-flight mass spectrometer (MAXIS; Bruker Daltonics).

2.12 | Cell and culture method

Three different CHO-K1-GS derived cell lines expressing different mAbs (mAb2, mAb3, mAb4) were thawed with a ThawSTAR automated cell thawing system (MedCision Inc). Precultures were cultivated in shake flasks of adequate size at 36.5°C, 5% CO2, and 120 rpm in Multitron incubators. Cultures were subcultivated every two or three days by diluting an appropriate volume of cell culture liquid with fresh medium in a new shake flask before the inoculation of the ambr15™ bioreactors.

2.13 | Bioreactors (ambr15)

Cells were cultivated in the Advanced Microscale Bioreactor System (ambr15; Sartorius AG) in a 14-day fed-batch culture. The seed density for all cell lines was 0.7E6 cells/ml. The cell culture conditions were maintained at a temperature of 34.5°C, DO setpoint of 50%, pH of 6.95 ± 0.25 and an agitation speed of 1000 rpm over the complete cultivation time. Total and viable cell numbers were monitored by a Cedex analyzer (Roche Innovatis GmbH), while metabolic parameters such as glucose, lactate, and ammonium were measured by the Konelab. For harvest, the cell culture broth was centrifuged at 7100g and 4°C for 15 min in a KR4i centrifuge (Thermo Fisher Scientific Inc). The supernatant was filtered through a syringe (BD Plastipak 20 ml; BD) using 0.7 µm prefilters (Ministart NML GF hydrophilic; Satorius Stedim Biotech GmbH) and 0.22 µm main filters (Ministart Ophthalsart; Satorius Stedim Biotech GmbH) and stored at −70°C until further processing. The redox potential was measured after harvest and filtration by using the EasyFerm Plus ORP Arc 120 (Hamilton) probe according to manufacturer’s description.

Cultivation experiments testing 0 g/L and 3 g/L cysteine in the process media were fermented in duplicates. The duplicates of mAb4 were pooled prior charge variant analysis. The effect of cysteine concentrations in the process medium and boli was conducted using an I-optimal DoE design. All models were checked to contain no aliasing for discussed effects, for consistency and statistical power.

3 | RESULTS

3.1 | Heterogeneous charge does not impact biophysical mAb1 structure and stability

The main focus of this study was to identify the root cause of acidic charge variants of an IgG1 mAb (mAb1). When separated by analytical WCX, 25% of acidic variants were observed for mAb1. The APG consisted of three overlapping peaks (Figure 2a). To understand the root cause of the acidic variant formation, mAb1 was fractionated to yield eight APG fractions (A–H), one main peak fraction (I) and two BPG (BPC; J–K) fractions (Figure S1). These sequential small volume fractions were collected to stepwise characterize the full APG spectrum. A control sample was prepared by pooling all fractions of
an individual preparative run. The purity of each fraction was confirmed by reinjecting the fractions onto the WCX column, immediately after buffer exchange (Figure 2a), and after 1 week of storage at 4°C (Figure S2). The charge variants were characterized for their biophysical and structural properties, as well as stability. Analytical SEC demonstrated a rather uniform distribution of LMW species (Figure 2b; Table S1). LMWs observed by analytical SEC originate from backbone fragmentation. Merely the most acidic fraction A contains slightly elevated levels of LMWs, which is expected from the literature (Du et al., 2012). HMW species, on the other hand, were more dominant in the most basic species K. The structural integrity of the charge variants was analyzed by far UV CD spectroscopy (Figure 2c). The minima at 219–220 nm and maxima at 202 nm observed for all samples were indicative for dominant β-sheet structural elements, and local minima at 209–210 nm implied α-helical contributions, as would be expected for a mAb (Buijs et al., 1996). The distinct overlap between the spectra suggested similar secondary structural elements and thus highly comparable structures in the individual charge fractions. The stability of the charge variants was assessed by nano-DSF, which revealed changes in intrinsic Trp fluorescence upon thermal unfolding (Figure 2d; Table S1). Two melting temperatures (T_m) were detected, corresponding to unfolding events of individual mAb domains, as well as the onset temperature for scattering, indicating aggregation.
CH2 and Fab domains unfolded at 69.4–70.0°C and 74.9–75.1°C, respectively. Overall, no differences in thermal unfolding or aggregation onset were found between the fractions, implying that charge heterogeneity does not affect the stability of mAb1.

3.2 Modified disulfide bonds contribute to acidic antibody species

To investigate the nature of the acidic variants, the individual fractions were analyzed by nonreducing CE, which allows a characterization of antibody integrity. Noncomplete mAb species originating from broken interchain disulfide bonds migrate as separate peaks on the CE under nonreducing (native) conditions. These species include the free LC, free HC, half-molecules (HL), two interconnected HCs lacking LCs (HH), and mAbs missing one LC (2H1L). In contrast to analytical SEC where these species would elute with the main peak, nonreducing CE is a valuable technique to evaluate the degradation of interchain disulfide bonds. Nonreducing CE demonstrated an increase of noncomplete mAbs in the more acidic fractions (Figure 3a; Table S1). In particular, the LC and 2H1L species were accumulated in early eluting acidic fractions (up to 8.2% of LC), showing increased interchain disulfide bond rupture in acidic fractions.

**FIGURE 3** Analysis of monoclonal antibody 1 (mAb1) cysteines revealed that rupture of intra- and intermolecular mAb1 disulfide bonds contribute to acidic charge variants. (a) The mAb1 charge variants were analyzed by nonreducing capillary electrophoresis (CE) to identify mAb1 size variants resulting from interchain disulfide bond degradation (n = 2). Full CE data for all size variants is shown in Table S1. (b–d) The mAb1 charge variants were further analyzed by a liquid chromatography–mass spectrometry (LC-MS) peptide mapping approach to quantify modifications of different mAb1 cysteine residues (n = 1). (b) LC-MS measurement of mAb1 charge variants was performed to quantify the relative levels of glutathionylations, free thiols, trisulfides, and cysteinylations for the Cγ-3 residue, which is the cysteine residue in the light chain typically forming an interchain disulfide bond to the heavy chain. (c,d) Using LC-MS peptide mapping, mAb1 cysteines typically involved in intrachain disulfide bonds in native mAb1 were analyzed. The levels of free thiol content (c) and cysteinylation (d) were quantified to investigate rupture and modification on intramolecular disulfide bonds. The cysteinylation content in Cγ3-1 in the mAb1 control is presumably an outlier. In addition, the levels of cysteinylation are shown for Cγ-3 (d) [Color figure can be viewed at wileyonlinelibrary.com]
Peptide mapping LC-MS measurements revealed that modifications of the C-terminal cysteine (C₃-3) of the LC are responsible for the abundance of free LC found in acidic fractions (Figure 3b). In the most acidic fraction APG A, relative levels of glutathionylations (9.3%), free thiols (4.8%), trisulfide bonds (0.67%), and also cysteinylations (0.05%), although to a lower extent, were found. In contrast to glutathionylations, free thiols and cysteinylations, which accumulate in acidic fractions, trisulfide bonds showed a constant distribution in all fractions, and therefore do not seem to contribute to acidic elution.

Next, the status of additional cysteines forming intramolecular bridges, which cannot be observed by CE, was investigated in the different mAb1 charge variants. In the peptide-map LC-MS approach, it was possible to analyze further mAb1 cysteines, which showed sufficient chromatographic separation and mass spectrometric properties. Regarding free thiols, the cysteines C₁₂-1, C₁₂-1, and C₁₂-2 (see Figure 3 for reference) could be analyzed and the experiment revealed, that the number of unpaired cysteines was increased in APG fractions with respect to the main peak and fractions from the BPG (Figure 3c). Intriguingly, the most acidic eluting APG A fraction contained the highest number of free thiols. With respect to cysteinylation, further three cysteines showed a comparable trend, namely cysteinyl in the APG, although at low absolute levels (Figure 3d). These investigated cysteines included two residues typically forming intrachain disulfide bonds in the HC (C₃-2-1 and C₃-3-1). Additionally, the cysteine involved in the interchain disulfide bond between the LC and 2H1L (C₂-3, also shown in Figure 3b) is shown for comparison.

In line with C₂-3 (Figure 3b), the presented data demonstrated that other cysteines are affected by modifications, although at lower levels. Taken together, these data strongly support that disulfide bond rupture and possible subsequent modification contribute to acidic variants. Yet, disulfide scrambling, another typical disulfide bond modification which can contribute to acidic variant formation, was not detectable by LC-MS (Table S2). In addition to the redox-sensitive modifications well known posttranslational modifications were found including N-acetylleuraminic acid, high mannose, lysine glycation, and N-terminal pyroglutamate formation (Figure S3). These modifications were distributed throughout the charge variants as expected from the literature (Chung et al., 2018; Du et al., 2012; Liu et al., 2008).

The peptide-mapping results indicate that redox-sensitive modifications are distributed throughout the mAb1 structure, however, only individual cysteines had been analyzed and most cysteines were not visible in this approach. To further locate the modification-sites, IdeS digestion of the charge variants was carried out, and the resulting F(ab)₂ and Fc fragments were subjected to intact LC-MS measurements. Heterogeneous peaks and degradation were observed for the F(ab)₂ fragment (Figure 4b), while the Fc fragment appeared as a homogeneous peak (Figure 4a), pinpointing the majority of cysteinylations and glutathionylations, as well as glycations, to the F(ab)₂ part of mAb1. It should be noted that this experiment can provide only a qualitative trend and no quantitative information for all cysteines.

3.3 | In vitro cysteine treatment induces acidic variants by breaking disulfide bonds and introducing cysteinylations

To prove that disulfide bond breakage contributes to acidic mAb1 species, in vitro experiments were conducted by adding cysteine to mAb1 and analyzing the resulting charge species (Figure 4). Cysteine treatment did promote increased levels of acidic peaks in mAb1, in a concentration-dependent manner (Figure 4a). Incubation with cysteine induced a uniform increase in APG levels, rather than enhancing particular peaks. BPG levels did slightly decrease, as well as the Main species, indicating that both were converted into more acidic species. In control experiments, intact mAb1 and the Main I species were incubated with S-sulfoceysteine and cysteine, nonreducing derivatives of cysteine (Figure S5). In contrast to cysteine, the control substances did not trigger changes in charge distribution, as shown by the WCX profiles.

These results indicate that incubation with cysteine is sufficient to generate the full APG profile of mAb1. This observation is further supported by an experiment, in which cysteine was added to the purified main species (referred to as Main I), which was free from acidic variants prior incubation. Remarkably, cysteine treatment caused the formation of multiple acidic species representing the full APG profile (Figure 4b). Analogs to cysteine, glutathione treatment caused a change in the mAb1 charge pattern, as illustrated by WCX (Figure 4c). However, while cysteine induced a uniform increase of the acidic peak group, glutathione targeted predominantly one specific signal, which enhanced the particular variant eluting at 18.5 min.

As expected, the reducing characteristic of cysteine led to the generation of mAb1 species eluting more acidic on the WCX. The resulting species of the non-fractionated mAb1 incubation with cysteine were further characterized by CE analysis, which demonstrated an increase of numerous variants, such as LC, HC, HH, and 2H1L after cysteine treatment of the nonfractionated mAb1 (Figure 4d). This data further supports that native mAb1 contains significant amounts of variants, which are the result of intermolecular disulfide bond reduction. The same trend was observed for the purified mAb1 Main I peak after cysteine incubation (Figure 4e). Cysteine concentrations of 1 mM triggered a more significant increase in APG levels in the Main I sample (3.4% to 14.4%) as compared to the nonfractionated mAb1 (25.1% to 28.9%). This result supports that in nonfractionated mAb1, the existing acidic variants are, to some extent, already composed of species induced by disulfide bond rupture. Thus, it required higher concentrations of cysteine (10 mM) to induce considerable disulfide bond breakage and modification to observe changes in the APG profile of nonfractionated mAb1, in comparison to the Main I species, where these pre-existent species were not present.

An LC-MS peptide-mapping analysis was performed to study the type of modifications introduced by in vitro incubation with cysteine to additional mAb1 cysteine residues. Several mAb1 cysteines showed sufficient chromatographic separation and mass spectrometric properties for analysis. The analyzed mAb1 cysteine residues...
included those involved in intramolecular disulfide bonds, which did not appear in CE analysis. Free thiol levels were slightly increased by incubation with cysteine for three intramolecular bonded cysteines (CH2-1, CL-1, CL-2; see Figure 1 for reference) (Figure 5a), thus showing the same trend as observed for intermolecular cysteines by CE (Figure 4d). In addition to the reduction to free thiols, intramolecular disulfide bonds of further cysteines (C1,2-1, C1,2-1, and C1,3-1) were susceptible to cysteinylation, in particular at high cysteine concentrations (Figure 5b). The same trends were observed for the interchain disulfide bonded C1,3-1 residue.

FIGURE 4  In vitro treatment of monoclonal antibody 1 (mAb1) with cysteines causes degradation of intermolecular mAb1 disulfide bonds. (a) Nonfractionated, intact mAb1 was treated with increasing cysteine concentrations (n = 4) for 2 h and subsequently analyzed by analytical weak cation exchange chromatography (WCX) to track changes in the charge variant profile. (b) Likewise, the fractionated, isolated mAb1 fraction Main I was treated with increasing cysteine concentrations for 2 h and analyzed by WCX to study the impact of cysteine incubation on acidic peak group formation (n = 4). As a control, the nonfractionated mAb1 is shown in gray. (c) WXC was carried out for mAb1 upon incubation with glutathione (GSH:GSSG (1:1)) for 0, 4, and 24 h (n = 2). (d,e) Nonreducing capillary electrophoresis was conducted after a 2 h incubation with increasing cysteine concentrations to observe the effect of cysteine on the rupture of intermolecular disulfide bonds (n = 4). This experiment was carried out for both the nonfractionated, intact mAb1 (d), as well as on the fractionated, isolated mAb1 fraction Main I (e) [Color figure can be viewed at wileyonlinelibrary.com]
3.4 | Cysteine changes the redox potential during fermentation and shapes mAb charge profile

Cysteine is an important amino acid and required for stabilizing cell viability in long-term cultivations as well as in elevating final product concentrations (data not shown). In previous experiments of this study, it had been shown that cysteine induces acidic mAb variants in vitro. To study the effect of cysteine during cell cultivation different amounts of cysteine were added to fermentation of three different mAbs (Figure 6). Usually cysteine is part of the production medium concentrations (data not shown). In previous experiments of this study, it had been shown that cysteine induces acidic mAb variants in vitro. To study the effect of cysteine during cell cultivation different amounts of cysteine were added to fermentation of three different mAbs (Figure 6). Usually cysteine is part of the production medium

FIGURE 5 Intramolecular monoclonal antibody 1 (mAb1) disulfide bonds are disintegrated and modified by in vitro cysteine incubation. Liquid chromatography–mass spectrometry peptide-mapping was conducted on mAb1 treated with increasing cysteine concentrations for 2 h. The peptide mapping approach allowed to study the state of several cysteines typically involved in intramolecular disulfide bonds in native mAb1. The levels of free thiols (a) and cysteinylation (b) were quantified to investigate rupture and modification on intramolecular disulfide bonds induced by incubation with cysteine (n = 1). In addition, levels were detected for C2-3, a cysteine residue involved in an interchain disulfide bond in native mAb1.

FIGURE 6 During fermentation, the media component cysteine can influence the redox potential and impact the resulting monoclonal antibody (mAb) charge profile. (a) Three mAbs were cultivated at 0 and 3 g/L cysteine concentrations in the process media (n = 2), and the acidic peak group (APG) levels, as well as the redox potential (plotted on the right y-axis), were detected at the end of the fermentations (mAb4 was pooled prior charge variant analysis). (b–d) DoE of three mAbs and the resulting APG profiles after addition of cysteine at different stages of fermentation. The model function of the main effects are depicted including the 95% confidence intervals (dotted lines). (b) APG levels as a result of different cysteine concentrations in the process medium (c) APG levels as a result of different cysteine concentrations in the bolus addition (d) APG levels as a result of the day of cysteine-bolus addition [Color figure can be viewed at wileyonlinelibrary.com]
and the feed medium, but it can also be given as an independent bolus during cultivation. When adding cysteine to the production medium, a trend towards lower redox potentials (favoring a reducing environment) was observed after 14 days of cultivation (Figure 6a). Simultaneously, APG levels increased in a molecule-independent manner. For mAb2 and mAb3, enhanced APG levels correlated with increased cysteine concentrations and lower redox potentials.

Varying cysteine concentrations in the production medium and bolus additions on different cultivation days were investigated (Figure 6b–d). While increased levels of cysteine in the production medium only contributed to a moderate elevation of APG levels (Figure 6b), large bolus addition of cysteine had a more pronounced effect on APG generation (Figure 6c). Acidic species increased at low cysteine concentrations added as bolus to the cultivation and eventually reached a plateau at higher cysteine levels. This effect was observed for all mAbs, although to different extents. Intriguingly, in contrast to concentration, the time point of bolus addition did not contribute to charge variant generation. Thus, addition at any point during fermentation resulted in full APG manifestation (Figure 6d).

4 | DISCUSSION

4.1 | Redox-sensitive modifications are a common feature of acidic variants of mAb1

A defective disulfide bond network has been reported to introduce different charge to mAbs (Bloom et al., 1997; Gevondyan et al., 2006; Gu et al., 2010; Schauenstein et al., 1986; Seibel et al., 2017; Wang et al., 2018). Although mAb charge variants have been characterized and described in the literature, insights to the underlying mechanisms of their generation would be beneficial in controlling mAb charge variants in industrial production processes. The current study provides an in-depth investigation of the origin, impact and control of acidic species of an IgG1 (mAb1). When separated by WCX, the APG level accounts for 25% of the total mAb1 charge population (Figure 2a). In the current study, eight mAb1 charge variants were separated, collected, and analyzed for their biophysical properties and modifications. Despite the different charge, the mAb1 species were comparable in terms of stability and structural integrity, as demonstrated by thermal unfolding and CD spectroscopy (Figure 2c,d). This data implies that mAb1 modifications leading to changes in charge do not affect stability and thus may be irrelevant for drug safety and efficacy. These findings are in contrast to an earlier study, where high levels of acidic species were associated with changes in the biophysical properties and bioactivity of an IgG (Banks et al., 2008). In the earlier report, the origin of these changes was traced back to cysteinylation of an unpaired cysteine in the complementary-determining region (CDR). It should be noted, that the investigated mAb1 in the current study does not contain cysteine residues in the CDR. Thus, the location of the affected cysteine, the availability of partner-cysteine for pairing, and the degree of modification seem to determine the criticality towards structure, stability, and activity.

Further investigation of the isolated acidic species by nonreducing CE and LC-MS revealed that degraded and modified disulfide bonds contribute considerably to the acidic species found in mAb1 (Figure 3). Noncomplete antibodies (e.g., missing LCs) were observed by nonreducing CE for acidic variants (Figure 3a) but not by analytical SEC (Figure 2b), thus, the noncomplete antibodies were most likely a result of degradation of disulfide bonds, leading to chain dissolution in nonreducing CE. The reduction of an interchain disulfide bond would not cause the LC to elute separately from the remaining mAb under the native conditions of SEC measurements, but the presence of SDS in the nonreducing CE measurement leads to separate migration of the LC.

Cysteine residues involved in inter- and intrachain disulfide bonds were modified in acidic mAb1 species, as shown by LC-MS (Figure 3b–d). The C-terminal cysteine of the LC which forms a disulfide bond to the HC has been described as being vulnerable to modifications (Wang et al., 2018). In acidic species, this cysteine contained considerable modifications, as compared to more basic species. These modifications included the reduction to free thiols, as well as cysteinylations and glutathionylations (Figure 3b).

4.2 | Modified cysteine disulfide bonds are distributed throughout the mAb structure

As revealed by a peptide-mapping LC-MS approach, affected cysteine residues in acidic species seem to be distributed to all mAb1 domains and include intra- and interchain disulfide bonded cysteines (Figure 3b–d). Detailed site-specific relative quantification by LC-MS was not feasible for all cysteine residues due to poor resolution and mass spectrometric behavior. Yet, a significant amount of redox-sensitive modifications contributing to acidic character originated from cysteines located in the F(ab)′2 region of mAb1, as demonstrated by intact LC-MS measurements of nonreduced IdeS-digested mAb1 (Figure S4). This is in line with earlier reports stating that IgG1s feature unpaired cysteines in their variable domains (Chumsae et al., 2009; Xiang et al., 2009). Results of the current study suggest that such cysteines are susceptible to cysteinylation and glutathionylation, in addition to being in a reduced oxidation state.

4.3 | The media component cysteine induces and enhances the full acidic variant profile

Cysteines of an intact intrachain disulfide bond in a natively folded antibody are more protected from degradation and modification, as compared to those involved interchain disulfide bonds (Amzel & Poljak, 1979; Kikuchi et al., 1986). The occurrence of nonintact
intrachain disulfide bonds in mAb1 indicates that the disulfide network is not properly formed during the production process. Cysteine as a media component contributes to the redox-environment during fermentation, which is critical for a correctly formed disulfide network (Chaderjian et al., 2005; Handlogten et al., 2017). As cysteinylations was a modification detected by LC-MS, the influence of cysteine on mAb1 was simulated in in vitro experiments. Indeed, the addition of cysteine alone was able to increase the number of acidic species (Figure 4a) concomitantly with disulfide bond degradation (Figure 4d) and residue-specific cysteinylation (Figure 5b), providing a strong link between disulfide bond rupture and the incidence of acidic species.

Cysteine incubation of the purified main peak of mAb1 strikingly resulted in the reproduction of the entire APG profile on a WCX column, featuring all peaks found in the control sample (Figure 4b). Thus, multiple acidic variants can be generated from "from scratch". The generation of all three major acidic peaks via in vitro incubation with cysteine implies that analogously, the media component cysteine may contribute considerably to mAb AGP levels during cell culture, in combination with further complex factors described previously (Chaderjian et al., 2005; Handlogten et al., 2017). Further, the findings provide insight into the mechanism of cysteine-induced APG formation. Modifications introduced by cysteine include cysteinylations as well as reduction of cysteines to free thiols resulting in non-complete antibodies. Thus, the WCX profile of native mAb1 must consist of various variants caused by disulfide bond degradation and potential subsequent modification. Further possible effects such as disulfide bond shuffling and mispairings were not detected for mAb1 (Table S2). Yet, posttranslational modifications also contributed to mAb1 acidic species formation, as seen from LC-MS (Figure S3) and as expected from the literature (Harris et al., 2001; Khawli et al., 2010; Lyubarskaya et al., 2006; Miller et al., 2011; Moorhouse et al., 1997; Yan et al., 2009), but did not explain the entire panel of acidic species observed.

Incubation of mAb1 with glutathione resulted in the increase of specific acidic peaks (Figure 4c), instead of an uniform increase of the entire APG, as in the case of cysteine. Thus, glutathione and cysteines may operate with different specificities. While the cysteine-induced mAb1 species were heterogeneous and characteristic for the full APG profile, glutathione seemed to generate a specific species, presumably a glutathionylated mAb1 LC. This assumption is supported by the observation of high degrees of glutathionylation found in the LC (Figure 3b). Furthermore, glutathione has been reported to disintegrate interchain disulfide bonds during fermentation (Handlogten et al., 2017). In comparison to glutathione, the mechanism of cysteine during in vitro treatment is less specific, as supported by the higher number of cysteine residues affected by cysteinylations (Figure 5b). Hypothetically, glutathione targets Cys-3 specifically, due to its solvent exposed location at the C-terminus of the LC. Steric hindrance may prevent glutathione from reacting with more solvent-protected cysteines. Furthermore, cysteine is often ranked as a hydrophobic residue (Poole, 2015), and may exert higher binding affinities to free cysteine, as they share the same hydrophobic properties, in comparison to glutathione. This may serve as an additional explanation why a higher number of mAb1 cysteines are modified by free cysteine than free glutathione.

As cysteine contributed significantly to the APG profile in vitro, its influence was investigated in fermentation experiments. Indeed, cultivation at increased cysteine concentrations in the process media enhanced the APG levels of three different mAbs (Figure 6a,b). A product-specific contribution of this effect may need to be taken into account, as not all investigated mAbs were affected to the same extent. In addition, APG generation was found to be dependent on cysteine concentration, as well as on the route of cysteine addition (i.e., bolus addition had the most impact; Figure 6c,d). Based on the in vitro data, it can be assumed that cysteine causes the rupture of disulfide bonds and introduction of cysteinylations during fermentation. The formation of trisulfide bonds has previously been linked to cysteine in the feed (Kshirsagar et al., 2012), and may additionally contribute to acidic species.

The media component cysteine is subjected to complex sulfur metabolism during cell culture (Ali et al., 2019). Cysteine has been shown to oxidize to cystine over 28 days in chemically defined media, yet 20% of the originally added substance remains in its reduced cysteine state (Krattenmacher et al., 2019). Thus, although it is not expected that the added absolute cysteine concentrations during cell culture reflect the levels present throughout fermentation, it can be assumed that the relative levels correlate. Cysteine therefore exerts a concentration-dependent effect on acidic species, which is highly relevant from an applied, bioprocessing perspective. An impact of cysteine during cell culture is not expected, based on in vitro experiments (Figure S5).

4.4 | Formation of acidic variants is most likely driven by redox reactions induced by cysteine in fermentation media

The modification of mAb disulfide bonds by cysteine may be driven by two possible mechanisms (Figure 7): the oxidized disulfide bond can undergo a Sπ2 substitution resulting in cysteinylations (1), alternatively, the cysteines of an oxidized disulfide bond will first be reduced to free thiols by a favorable redox potential and subsequently be cysteinylated by excess cysteine in solution (2). Numerous results from this study support the occurrence of free thiols and highlight the significance of the right redox potential for an intact disulfide bond network, thereby supporting the second scenario. Firstly, S-sulfocysteine and cystine, nonredox active derivatives of cysteine, were not able to alter mAb1 WCX profiles (Figure S5). Secondly, free thiols were detected in highly acidic mAb1 species (Figure 3b,c). Thirdly, in cultivation experiments of mAb2 and mAb3, the addition of cysteine to the production medium was not only associated with an increase in acidic variants, but also a decrease in redox potential, favorable for disulfide bond reduction (Figure 6a). Although Sπ2 substitution by cysteine in the media may occur, the mechanism via redox potential is more likely, as it contributes to acidic species associated with defective disulfide bonds.
FIGURE 7 Possible mechanisms of cysteine-induced cysteinylation of monoclonal antibody disulfide bonds. (1) $S_n^2$ substitution of intact disulfide bonds resulting in cysteinylation. (2) Redox reaction leads to a reduction of disulfide bonds followed by cysteinylation of the free thiols. Results of the study point to the second scenario as the most likely mechanism.

The findings reported in this study illustrate the APG-promoting effect of cysteine in cell culture media for an IgG1 resulting in disulfide bond degradation and modification. Elevated levels of acidic species are undesirable during bioprocessing as they impair step yields. The demonstrated data highlights cysteine concentration in cell culture media as a potential point of interference for manipulating mAb charge during the production process. Specifically, careful monitoring of cysteine as a media component and thus regulation of the redox potential during fermentation may enable a control of charge heterogeneity.

ACKNOWLEDGMENTS

Boehringer Ingelheim Pharma GmbH & Co. KG. funded all research of this manuscript. The authors are especially grateful to Göran Hübner, Nils Drechsler and Sarah Lergenmüller for scientific discussions and input on analytical data, as well as Oliver Burkert for providing the opportunity for circular dichroism spectroscopic measurements. Further, the authors would like to thank Michelle Ahlers-Hesse for technical support in purification and in vitro experiments, Elena Bollgoen and Chris Schubert for their contributions in the cell culture experiments, as well as Amadine Calvet for measurements of redox potentials.

AUTHOR CONTRIBUTIONS

Elke Prade and Ingo H. Gorr designed the research and wrote the manuscript with input from all authors. Anne Zeck measured and analyzed the liquid chromatography–mass spectrometry data. Fabian Stiefel and Andreas Unsoeld designed and analyzed the cell culture experiments. David Mentrup performed the sample preparation and analytical characterization of fractionated and intact mAb1 and analyzed the data. Elke Prade and Erik Arango Gutierrez supervised the mAb1 wet lab activities. All authors discussed the results.

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**How to cite this article:** Prade E, Zeck A, Stiefel F, et al. Cysteine in cell culture media induces acidic IgG1 species by disrupting the disulfide bond network. *Biotechnology and Bioengineering, 2021;118:1091–1104.*

https://doi.org/10.1002/bit.27628