The Effect of Gadolinium on the Microstructures and Charpy Impact Properties of Super Duplex Stainless Steels

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Abstract: Super duplex stainless steels (SDSSs), exhibiting excellent strength and corrosion resistance, serve as the attractive materials in a variety of industries. However, improvements in their ductility and impact-toughness are required in extreme environments. In this study, the effects of gadolinium on the microstructures and Charpy impact properties of super duplex stainless steels were investigated. A base super duplex stainless steel (BDSS) and a gadolinium-added super duplex stainless steel (GDSS) were successfully fabricated using an air casting method. The oxygen content and grain size of SDSSs were found to decrease because of high reactivity of gadolinium with oxygen. Moreover, the average inclusion size and area of GDSS also decreased even with a slight decrease in the average distance between inclusions. Both the BDSS and GDSS exhibited typical Charpy impact transition behavior from −196 °C to 200 °C. Moreover, the GDSS impact energies using Charpy test were higher than those of BDSS over the entire temperature range. Moreover, the ductile-to-brittle transition temperature (DBTT) of SDSSs calculated from the fracture appearance transition temperature (FATT) significantly decreased by over 20 °C with the addition of gadolinium.

Keywords: duplex stainless steel; gadolinium; Charpy impact properties; microstructure; inclusions; grain size

1. Introduction

Recently, rare-earth metals (REMs) have been widely used in various materials to improve their performance or change their functionality [1]. A small addition of REMs in the materials can cause important changes in the properties of the materials. REM (such as, ceria and lanthana)-based compounds are receiving increasing attention due to their automotive exhaust catalyst applications, high oxygen storage capacity and larger ionic size [1–3], which result in a significant enhancement of the catalyst performances. A major application of REMs is remarked in the luminescent materials, such as phosphors [4–7]. REMs can also be used as additives in cast metals to improve their microstructures and mechanical properties. However, when they react with harmful non-metallic inclusion-forming elements, such as nitrogen, oxygen, and sulfur, they can modify MnS or MnO, resulting in the formation of stable REM compounds, such as REMO_{x}S_{y} [8–11]. Some researchers reported that the addition of REMs improved the mechanical properties of iron and steel by changing the graphite and sulfide morphologies, respectively [12–15]. Additionally, REMs can reduce the grain size of cast metals,
influencing the strength and toughness of the metals [9,16]. Using REMs for various applications in the plant, automotive, and aerospace industries, attempts have been made to fabricate high-quality cast metals.

Gadolinium (Gd), a specific type of REM with high neutron-absorbing ability, is commonly used in medicine to target tumors. Recent studies have also demonstrated its applications as a neutron-absorbing material which can be used in shielding of nuclear fuels in nuclear reactors [1,17]. Some researchers also identified Gd as a metal additive. Using a metal injection molding (MIM) method, Holm et al. fabricated Ti–6Al–4V alloys with 1–5 wt. % Gd [18]. The ability of Gd to reduce the grain size, change the residual pore size and shape after sintering enhances fatigue behavior. Ghandvar et al. fabricated an Al-15%Mg2Si in situ composite by adding Gd [19]. Typically, Gd was an important component that caused the formation of small Al3Gd particles, which acted as the heterogeneous nucleation substrates for Mg2Si, thereby modifying the composite’s macrostructure. Despite the outcomes of these pieces of research, only a limited number of studies investigated Gd as an additive use. Therefore, there is a need for more in-depth studies to understand the effect of Gd as a REM additive.

Stainless steels are one of the most commonly used metals across industries. Many different types of stainless steel are classified by their main phase and characterized by their various applications [20–22]. Ferritic stainless steels with high chromium, and low carbon contents exhibit good corrosion resistance and stress corrosion cracking with relatively low ductility [23–25]. Austenitic stainless steels, referred as non-magnetic steels, are characterized with good workability, high toughness, weldability and plasticity. However, they are vulnerable to localized corrosion in chloride-rich environments [26–28]. On the other hand, duplex stainless steels, comprising both ferrite and austenite phases, have received a great attention owing to their excellent corrosion resistance and mechanical properties [29–31].

Super duplex stainless steels (SDSSs) are a family of duplex stainless steels (DSSs) with a high pitting resistance equivalent number (PREN) over 40 [32,33]. SDSSs have much higher corrosion resistance than other DSSs, and thus, they are widely used in severe industrial environments, such as in the petroleum, nuclear reactor, and sea-plant industries [21,34]. Thus, their mechanical properties in the actual environment are quite important to ascertain the reliability of SDSSs under high-and low-temperature conditions [35]. Particularly, the impact toughness of SDSSs under extreme temperatures is important for the safety and long-term reliability of cryogenic machinery [36–38].

The purpose of this research is to investigate the influence of Gd on the microstructures and Charpy impact properties of SDSS. A890 grade 7A super duplex stainless steel was used and 0.1 wt. % of Gd was added and formulated for casting. Moreover, electron backscatter diffraction (EBSD) and field-emission scanning electron microscopy (FE-SEM) were carried out to investigate the microstructures and inclusion of the fabricated alloys. The phases and compositions of the alloys were characterized by X-ray diffraction (XRD) and energy-dispersive X-ray spectroscopy (EDS). The mechanical properties were examined using Charpy impact tests carried out at a different of temperatures.

2. Materials and Methods

2.1. Materials and Alloy Fabrication

In this study, all the SDSSs used for the experimental purpose were cast in a high-frequency induction melting furnace (Inductotherm, Rancocas, NJ, USA). The starting materials for SDSS fabrication were high-purity iron (Fe), nickel (Ni), chromium (Cr), manganese (Mn), silicon (Si), ferromolybdenum (Fe-60 wt. % Mo), and ferrochromium nitride (Fe-60 wt. % Cr-10 wt. % N). All metals excluding Gd were melted at approximately 1640 °C. Moreover, 0.1 wt. % Gd granules (1–2 mm, 4N grade, Treibacher industrie AG, Althofen, Austria) were added to the molten SDSSs at approximately 1580 °C to suppress the oxidation of Gd. The air-cooled cast SDSSs in a sand mold were
cut into 200 mm × 30 mm × 30 mm specimens and solution-treated for 2 h at 1130 °C. The specimens were then quenched in water to avoid secondary phase formation and to achieve an approximate ferrite-to-austenite ratio of 50:50 according to the ASTM A890/A890M standard [39]. The chemical compositions of the fabricated alloys were analyzed using a gas analyzer (ONH-2000, Eltra, Haan, Germany) and inductively coupled plasma optical emission spectroscopy (ICP-OES) (iCAP-6300, Thermo Scientific, Waltham, MA, USA). The results are presented in Table 1. The alloys fabricated without the addition of Gd is referred as BDSS (base duplex stainless steel) while the alloy fabricated with the addition of Gd is called GDSS (Gd-added duplex stainless steel).

| Table 1. Chemical compositions of standard A890 grade 7A duplex stainless steel, BDSS, and GDSS (wt. %). |
|---------------------------------------------------------------|
|                  | Cr  | Ni  | Mo  | Mn  | W   | Cu  | Si  | N   | O   | Gd | Fe  |
| Standard         | 26–28 | 6–8 | 2–3.5 | 1–3 | 3–4 | −1  | −1  | 0.3–0.4 | −  | −  | Bal. |
| BDSS             | 27.3 | 6.8 | 3.7  | 1.1 | 3.4 | 0.7 | 0.3 | 0.3243 | 0.0692 | −  | Bal. |
| GDSS             | 26.9 | 6.9 | 3.1  | 1.2 | 3.3 | 0.7 | 0.4 | 0.3157 | 0.0568 | 0.016 | Bal. |

2.2. Microstructural Analysis

The microstructural characterization’s specimens were prepared by polishing the surface up to 2000 grit SiC paper and finishing using alumina suspension slurries. The microstructural analyses of the fabricated SDSSs were performed after solution treatment with an optical microscope (ECLIPSE MA200, NIKON, Tokyo, Japan) and the backscattered electron (BSE) detector of a Quanta 200F field-emission scanning electron microscope (FE-SEM; FEI Company, Hillsboro, OR, USA). In addition, electron backscatter diffraction (EBSD; FEI Company, Hillsboro, OR, USA) analysis was used to measure the grain size and ferrite-to-austenite ratio by assuming that Fe-BCC (Body centered cubic) for ferrite and Fe-FCC (Face centered cubic) for austenite exists in SDSSs. All EBSD data was obtained using the following parameters: working distance of 15 mm, accelerating voltage of 20 kV, 2.5 μm step size, and magnification of ×55. The measurement of an area of 4 × 4 mm² takes about 10 h. Orientation imaging microscopy analysis, provided by Explorer 4 Analyzer with Metals Quality Analyze (MQA) software, was used in conjunction with the FE-SEM. Three images of each SDSSs were examined to obtain the mean and standard deviation of the grain size and ferrite-to-austenite ratio. The average grain size of each SDSSs was calculated from the measured ratio of ferrite and austenite, and each grain size of ferrite and austenite. The ferrite content was additionally measured using a ferrite scope (Feritscope, FMP-30, Fischer, Achern, Germany) and a ferrite detection limit of 0.1%. Approximately 30 points on each specimen were analyzed. The chemical compositions of the specimens, while observing the microstructures, were analyzed using an energy dispersive X-ray spectroscopy (EDS) attached to the FE-SEM. Furthermore, the phases of the SDSSs were characterized by X-ray diffraction (XRD, D8-discover, Bruker, Germany) with monochromatic Cu Kα radiation over a diffraction range of 10–110° at a scanning rate of 0.1°/min.

The size distribution and average areas of the SDSSs inclusions were analyzed, using a CTR 6000 inclusion analyzer (Leica, Wetzlar, Germany) to analyze 30 points of each specimen. The average areas of the inclusions are reported as a mean ± standard deviation. One-way ANOVA was used to compare the means of the groups, and the level of significance was p < 0.05. The average distance between inclusions ($X_0$) in the fabricated SDSSs is defined as:

$$X_0 = 0.89R_0f^{-1/3}$$

where $R_0$ represents the average radius of the inclusions in the SDSSs and $f$ is the fraction of the inclusions [40,41]. It is assumed that the inclusions are spherical and distributed homogeneously in the SDSSs.
2.3. Charpy Impact Property Evaluation

The analysis of the mechanical properties of BDSS and GDSS was carried out using Charpy impact tests, performed following ASTM E23 on V-notched specimens (10 × 10 × 55 mm³) at temperatures between −196 °C and 200 °C, using a Charpy-type impact tester (SA-ID3, Satec, Union, NJ, USA) [42]. The Charpy analysis was performed on four specimens for each experimental SDSS group to obtain statistically significant results.

The fracture modes of the specimens were analyzed using FE-SEM and EDS. Based on the results from the Charpy impact transition curve, the lower shelf energy (LSE) and upper shelf energy (USE) were determined [43,44]. The ductile-to-brittle transition temperature (DBTT) was calculated with two methods: one was based on the energy transition temperature (ETT), which was the average value of LSE and USE. The other method was based on the fracture appearance transition temperature (FATT) at which the area fractions of the cleavage (brittle texture) to dimple (ductile texture) is 50:50 from the fractured surface of specimens [45,46]. The average fraction of shear fracture, obtained from the surface fracture of SDSSs, was analyzed with more than three FE-SEM images for each SDSS reported as mean ± standard deviation.

The values of impact transition temperature (ITT) were chosen as 27J-ITT and 68J-ITT (corresponding to Charpy impact energy absorption at 27J and 68J, respectively) following the Wallin master curve and ASTM standard in BS 7910:1999 and ASTM E185-82, respectively [47,48].

3. Results and Discussion

3.1. Microstructural Characterization

ASTM grade 7A SDSS alloys, which mainly comprised of 27% Cr, 7% Ni, 3% Mo, and 0.35% N, were successfully fabricated by casting. Both the fabricated BDSS and GDSS did not show any noticeable voids or cracks, and their chemical compositions matched the targeted composition (ASTM A890 grade 7A) as shown in Table 1 [39]. Moreover, the BDSS and GDSS’s PREN values, calculated with the following equation: PREN = wt. % Cr + 3.3 × (wt. % Mo + 0.5 × wt. % W) + 16 × wt. % N, were greater than 50, indicating an excellent pitting corrosion resistance [49]. The result suggests that both BDSS and GDSS can withstand extreme chemical conditions under the chloride, sulfide, or hypochlorite ion-containing solutions.

Upon addition of Gd, the oxygen content of SDSSs decreased from 692 ppmw to 568 ppmw, which was about 20% lower. Most research indicates that REMs can deoxidize cast metals, such as steels [50–52]. In this study, we demonstrated that Gd effectively reduced the oxygen content in SDSSs by reacting with oxygen during casting.

Figure 1A,B show the optical micrographs of as-cast SDSSs. Both types of SDSSs, regardless of their Gd’s contents, exhibited typical microstructures of duplex stainless steels. The light gray regions in the images correspond to the austenite phase (γ), and the dark gray regions correspond to the ferrite phase (α) matrices in BDSS and GDSS. No undesired secondary phases, such as sigma (σ) and chi (χ) phases were detected in both SDSSs, suggesting that the solution treatment at 1130 °C was effective. However, some irregularly shaped and heterogeneously sized spots were observed in BDSS, indicating the presence of non-metallic inclusions, whereas only a few smaller spots were observed in GDSS.

EBSD inverse pole figure maps of BDSS and GDSS are shown in Figure 1C,D, respectively. It should be noted that both EBSD images have random crystallographic texture owing to the absence of any plastic deformation. Furthermore, GDSS exhibited smaller grain size in ferrite region, whereas BDSS exhibited coarse grains. The calculated grain size of ferrite for BDSS and GDSS were 1026.8 ± 425.8 µm and 906.2 ± 297.6 µm, respectively. The results indicate that the grain morphologies of SDSSs are successfully modified with the addition of Gd without other phase changes. On the other hand, the grain size of austenite for both BDSS and GDSS were 178.4 ± 88.6 µm and 183.3 ± 107.8 µm and had no significant difference between them. Generally, it is well known that growth and recovery rates of austenite are slower than those of ferrite [53].
Figure 1. The microstructures of solution-treated duplex stainless steels (A) without gadolinium (BDSS) (B) with gadolinium (GDSS) and EBSD inverse pole figure maps of (C) BDSS and (D) GDSS.

The ferrite fraction and average grain size of SDSSs evaluated by EBSD mapping are shown in Figure 2. The ferrite contents in BDSS and GDSS were 57.7 ± 3.2% and 59.3 ± 2.4%, respectively, which were the appropriate ratios for duplex stainless steels, close to the calculated values of a ferrite scope. This result demonstrated that Gd did not affect the main phase of SDSSs. Meanwhile, the average grain size of SDSSs remarkably decreased from 683.5 μm to 598.3 μm after adding the Gd. During casting, Gd reduced the molten SDSSs surface tension thereby decreasing the activation energy of the nuclei formation [54]. An increase in several nuclei led to a decrease in grain size, which improved the mechanical properties of SDSSs owing to the increase in the number of grain boundaries, which interrupted dislocation movement in SDSSs [16].

Figure 3 shows X-ray diffraction patterns of BDSS and GDSS after solution treatment. The main peaks in the 2θ are ranged between 40° and 45°, corresponding to the (111) and (110) planes of austenite and ferrite, respectively. Only austenite and ferrite peaks are observed in both SDSS specimens. After solution treatment, the σ and χ phases that affected the steels mechanical properties negatively were removed. The result is in good agreement with the optical micrograph images (Figure 1). No differences in peak intensity between BDSS and GDSS were observed, suggesting that no unexpected reactions, such as SDSS’s phase transformation occurred after adding a small amount of Gd.
Figure 2. The grain size distributions of as-cast SDSSs for each phase characterized by electron backscatter diffraction (EBSD).

Figure 3. X-ray diffraction patterns of BDSS and GDSS.

3.2. Characterization of Inclusions

The microstructure changes of inclusions in SDSSs, casted with and without Gd, were further investigated by FE-SEM, as shown in Figure 4. Inclusions were detected in both ferrite and austenite phases. No clusters of inclusions were formed, and the inclusions were homogeneously distributed in both fabricated SDSSs (Figure 4A,B). Based on the BSE images polished surfaces, some irregular inclusions were found in BDSS, whereas relatively small inclusions were found in GDSS. The main
morphology of the inclusions in BDSS was polygonal and the size of the inclusions was around 20 µm, as shown in the high-magnification BSE image (Figure 4C). GDSS has two types of inclusions: polygonal inclusions as observed in BDSS and smaller spherical inclusions as shown in Figure 4B. In many previous reports, it has been proven that REMs can reform the inclusion shapes from angulated to non-angulated or spherical shapes with the ability to enhance the mechanical properties of metal alloys, such as the impact toughness [10]. The non-angulated inclusions in GDSS comprised of dark and light regions (Figure 4D). This type of inclusion was observed in steels modified by REM additives. These results suggest that the non-metallic inclusions detected in GDSS originate from the deoxidation reactions upon the addition of Gd [55].

![Backscattered electron micrographs of (A,C) BDSS and (B,D) GDSS. (A,B) were taken at low magnification and (C,D) at high magnification.](image)

The EDS was also used to quantify the inclusions’ chemical compositions, and the representative inclusions observed in both BDSS and GDSS (Figure 4C,D) were analyzed, as shown in Table 2. The observed elements contained in the inclusions in BDSS were O, Cr and Mn and their weight fractions were 21.8%, 50.9%, and 27.3%, respectively. Based on the EDS results, the polygonal inclusions in BDSS consist of Cr₂O₃ and MnO inclusions, which represent non-metallic inclusions formed in steels during casting [56,57]. The EDS analysis of inclusions in GDSS was performed for both the light regions and dark regions. Gd, Si, and Fe, as well as O, Cr, and Mn, which were the components of the inclusions in BDSS, were the main elements of the inclusions in GDSS. The EDS analysis results reveal that Gd-based oxides are the core of the complex inclusions, and (Mn, Cr, Si)-based oxides surround
the Gd-based oxides, such as Gd$_2$O$_3$. Thus, the addition of Gd to SDSS resulted in the transformation of the non-metallic inclusions in SDSS, as well as the size grains reduction. Moreover, the addition of a small amount of Gd with an undisclosed SDSS deoxidizing capacity resulted in a substantial change in the oxide content. Thermodynamically, REM-based oxides or oxysulfides are stable compounds owing to their formation of low standard free energies [58]. The standard Gibbs free energy formation for Gd$_2$O$_3$ is $-2457$ kJ/mol is larger than those for MnO, Cr$_2$O$_3$, and SiO$_2$. Therefore, it is expected that Gd-based oxides were formed in the melting state, followed by reaction of the remaining O with the molten SDSS and other elements, such as Si, Mn, and Cr.

**Table 2.** Quantitative chemical composition of the inclusions in each BDSS characterized by EDS (wt. %).

| Specimen  | O   | Cr  | Mn  | Fe  | Si  | Gd  |
|-----------|-----|-----|-----|-----|-----|-----|
| BDSS      | 21.8| 50.9| 27.3| -   | -   | -   |
| GDSS (light) | 15.0| 10.1| 9.9 | 7.3 | 10.1| 47.6|
| GDSS (dark) | 19.2| 41.1| 23.6| 12.4| 3.7 | -   |

Figure 5 presents the inclusion size distributions of the as-cast SDSSs and the average area of the inclusions, as well as the average distance between the inclusions in BDSS and GDSS. As depicted in Figure 5A, the inclusion size distribution shifts slightly to the left, indicating a decrease in inclusion size upon the addition of Gd. GDSS showed a bell-shaped curve, similar to a normal distribution centered at 6 µm, whereas BDSS had various sizes of inclusions with coarse inclusions over 20 µm. Approximately 85.2% and 92.9% of the inclusion sizes were between 0 and 11 µm in BDSS and GDSS, respectively. The calculated average size of the inclusions in GDSS was about 5.70 µm, which was an 18.7% reduction compared with the inclusions in BDSS (approximately 7.01 µm). This result demonstrated that the addition of Gd to SDSS significantly reduced the number of coarse inclusions and the average inclusion size.

**Figure 5.** (A) Inclusion size distributions of as-cast SDSSs. (B) the average area of inclusions and the average distance between inclusions in BDSS and GDSS (** $p < 0.005$).

The area of the inclusions per frame in GDSS significantly decreased compared with the area of the inclusions in BDSS, as shown in Figure 5B ($p < 0.005$). The decrease in the total area of inclusions was attributed to a decrease in the total O content with the addition of Gd. This result was consistent with the results observed in the optical image of BDSS, comprising of many heterogeneous inclusions (Figure 1A).

Rice et al. investigated the effects of inclusion fraction and the spacing between the inclusions on the ductile fracture behavior [59]. They developed a model called the Rice and Johnson model. In their model, the average nearest-neighbor spacing within the inclusions ($X_0$) is calculated with the equation given in Section 2 [59]. In this study, the distances between the inclusions in both BDSS and GDSS were quantified by calculating the sizes and fractions of the inclusions. As Gd was added to
SDSS, the average distance of the inclusions marginally decreased from 31.4 µm to 28.2 µm. With a simultaneous decrease in the inclusion size and fraction, the distances between inclusions were similar. The number of inclusions per unit area (\(N_A\)) is directly proportional to the fraction of inclusions and inversely proportional to the area of each inclusion:

\[N_A \propto \frac{f}{R_0^2}\]

From this equation, the number of inclusions per unit area in GDSS is larger than that of BDSS by about 11%. It should be noted a greater number of smaller inclusions were distributed in GDSS compared with BDSS.

3.3. Charpy Impact Tests of SDSSs

The influence of the inclusion structure and properties with the addition of Gd on impact toughness was investigated using Charpy impact energy tests. Figure 6A,B show the fracture specimens after Charpy V-notch tests at various temperatures from \(-196^\circ C\) to \(200^\circ C\) for BDSS and GDSS, respectively. As the experimental temperature decreased from \(200^\circ C\) to \(-196^\circ C\), the fracture surfaces of BDSS had a mirror-like shiny appearance (Figure 6A). Moreover, the surfaces of the specimens at lower temperatures were flat and the original shape was maintained, whereas the surfaces of the specimens at higher temperatures seemed to be bumpy with irregular shapes. The fracture surfaces exhibited both shiny regions and dull regions, corresponding to brittle and ductile fracture, respectively. These results suggested that a ductile-to-brittle transition occurred because of a change in experimental temperature. The GDSS fracture surfaces at various temperatures were similar to those of BDSS, as shown in Figure 6B. However, the fracture surface of GDSS at \(0^\circ C\) seemed to be completely rough, whereas that of BDSS at \(0^\circ C\) still had some shiny regions, indicating that the GDSS fracture mode changed at a lower temperature.

![Figure 6](image-url)

**Figure 6.** Fracture specimens after Charpy V-notch tests at various temperatures from \(-196^\circ C\) to \(200^\circ C\) for (A) BDSS and (B) GDSS.
Figure 7 shows the BDSS and GDSS Charpy impact energies from $-196^\circ C$ to $200^\circ C$. The Charpy impact tests were performed at ten different temperatures ($-196^\circ C$, $-100^\circ C$, $-60^\circ C$, $-40^\circ C$, $-20^\circ C$, $0^\circ C$, $25^\circ C$, $60^\circ C$, $100^\circ C$, and $200^\circ C$) to construct a sufficiently detailed absorbed energy curve that could be used to calculate various impact transition temperatures, as well as the ductile-to-brittle transition temperature (DBTT). The SDSSs impact energies as a function of experimental temperature can be expressed by a hyperbolic tangent function proposed by Oldfield as follows [60]:

$$\text{Charpy impact energy} = A + B \tanh \frac{T - T_0}{C}$$

where $T$ is the experimental temperature, $T_0$ is the mid-transition temperature, and $C$ is the slope of at $T_0$. Moreover, $A + B$ and $A - B$ are the upper shelf energy (USE) and lower shelf energy (LSE), respectively [60]. The constants $A$, $B$, $C$, and $T_0$ in this equation were calculated for both BDSS and GDSS, as shown in Table 3. Generally, specimens in the low temperature region have limited movement of domain walls and dislocations, whereas thermal activation permits more movement of domain walls and dislocations, resulting an increase in impact energy [61]. Charpy impact energy values for GDSS were larger than those of BDSS at all the experimental temperatures. The GDSS measured of LSE and USE were $8.0$ J and $169.8$ J, and those of BDSS were $6.0$ J and $156.0$ J, respectively. The theoretical LSE and USE for both SDSSs calculated from constants $A$ and $B$ were similar to the measured values, indicating that the obtained Charpy impact energy transition curves of both SDSSs were well-fitted by typical transition curves. In other words, three distinctive regions, namely lower shelf region, transition region, and upper shelf region, were observed. The mid-transition temperature shifted from $-32.3\, ^\circ C$ to $-37.9\, ^\circ C$ by the addition of Gd, corresponding to a decrease in DBTT by approximately $6\, ^\circ C$.

![Figure 7. Fracture Charpy impact energies of BDSS and GDSS as a function of temperature from $-196\, ^\circ C$ to $200\, ^\circ C$.](image)

**Table 3.** Impact of energy transition curve parameters and absorbed energy obtained from Charpy impact tests from the fabricated SDSSs.

| Specimen | $A$ | $B$ | $T_0$ | $C$ | Absorbed Energy (J) |
|----------|-----|-----|-------|-----|---------------------|
|          |     |     |       |     | LSE  | USE  |
| BDSS     | 80.2| 74.6| $-32.3$| 50.4| 6.0  | 156.0|
| GDSS     | 87.6| 78.6| $-37.9$| 43.5| 8.0  | 169.8|
The FE-SEM was used to examine the fracture surfaces of both SDSSs after the Charpy impact tests and the results were presented in Figure 8. The fracture surfaces of both BDSS and GDSS in the upper shelf region exhibited ductile fracture, as indicated by a variety of dimples (Figure 8E, J). Whereas cleavage with torn ridges indicating brittle fracture was observed in the lower shelf region (Figure 8A, F). However, a small increase in Charpy impact energy at cryogenic temperatures occurred upon the addition of Gd, whereas a larger difference between the Charpy impact energies was observed at high temperature as shown in Figure 7. At temperatures near the mid-transition temperature, the fracture modes for both SDSSs comprised both dimple and quasi-cleavage modes (Figure 8B–D, G–I).

Figure 8. Low-magnification FE-SEM images of fractured surfaces of BDSS (A–E) and GDSS (F–J) after Charpy impact tests at (A, F) −196 °C, (B, G) −60 °C, (C, H) −20 °C, (D, I) 50 °C, and (E, J) 200 °C.

Figure 9 shows high-magnification FE-SEM images of the fracture surfaces for both SDSSs at temperatures of −60 °C, −20 °C, and 50 °C analyzed to study the change in fracture mode in the transition region. The fracture surface of BDSS at −60 °C exhibited a cleavage-dominated fracture behavior with complex river patterns (Figure 9A). However, the fracture surface of GDSS at −60 °C, which had a 10% higher absorbed energy than BDSS, showed a greater proportion of various sizes of dimples and cleavage facets (Figure 9D). At −20 °C, the absorbed energy of BDSS increased by over 20% relative to that of GDSS and the absorbed energy values for BDSS and GDSS were 99.4 J and 122.5 J, respectively. The fraction of quasi-cleavage facets with small cleavage steps was approximately two thirds in BDSS (Figure 9B), and the ductile fracture areas covered approximately half of the whole fracture surface in GDSS (Figure 9E). On the other hand, at 50 °C, many dimples arising from ductile fracture appeared in both BDSS and GDSS (Figure 9C, F). However, most of the dimples in GDSS were bigger and deeper than those in BDSS. In general, the larger and deeper the dimple, the better plastic properties [62, 63]. These results are in accordance with a previous report that reveals the relationship between dimple structure and mechanical properties of steels [64]. Das et al. confirmed that the ductility value of a copper-strengthened high-strength low-alloy (HSLA) was directly proportional to the diameter of dimples.

In Table 4, various impact transition temperatures (ITT) calculated from Charpy impact tests are listed. According to BS 7910:1999, ASTM E185-82, 27J-ITT and 68J-ITT are generally used to evaluate the Charpy impact transition behavior of ferritic steels and nuclear reactor steels [47,48]. The temperatures at which the Charpy impact energy was 27 J (27J-ITT) for BDSS and GDSS were estimated at −77.3 °C and −82.4 °C, respectively allowed fitting of the data with the hyperbolic tangent function. When the impact toughness is 68 J (68J-ITT) for BDSS and GDSS, the temperature was −40.6 °C and −49.0 °C, respectively, around 35 °C higher than 27J-ITT. Remarkably, both 27J-ITT and 68J-ITT values decreased by 5–8 °C upon the addition of Gd, suggesting that the Charpy impact property was significantly affected by Gd addition.
Table 4. Impact transition temperatures calculated from Charpy impact tests on the SDSSs.

| Specimen | 27J-ITT (°C) | 68J-ITT (°C) | DBTT-ETT (°C) | DBTT-FATT (°C) |
|----------|--------------|--------------|---------------|---------------|
| BDSS     | 77.3         | 40.6         | 31.8          | 0.1           |
| GDSS     | 82.4         | 49.0         | 38.2          | 22.4          |

By determining the DBT Ts average value of LSE and USE (DBTT-ETT), the temperature was approximately 
−31.8 °C and −38.2 °C for BDSS and GDSS, respectively, and these values were quite similar to 
the mid-transition temperatures (Table 3). Therefore, significant improvements in Charpy impact 
properties (with USE of 169.8 J, 68J-ITT of −49 °C, and DBTT-ETT of −38.2 °C) were achieved by 
modifying the microstructure of SDSS with a very small input of Gd. The mechanism is explained as 
follows: First, Gd acts as a grain refiner, reducing the DBTT and increasing the USE. Second, the size 
and fraction of non-metallic inclusions are decreased by the reduction of O in SDSS with Gd acting as a 
deoxidizer. Although the number of inclusions increases and the average distance between inclusions 
decreases, the advantages of Gd addition make the modified SDSS a potential structural metal that can 
function under the extreme conditions. Moreover, the change in the shape of inclusions as globular 
shape might result in better impact toughness.

The DBTT can be evaluated with five methods: the energy transition temperature (ETT) which 
is considered to be the average value of LSE and USE, lateral expansion, shear fracture appearance, 
load diagram, and master curve [65–68]. In this study, an additional method based on fracture 
appearance transition temperature (FATT) was used. Figure 10 shows the average fraction of dimples 
obtained from surface fracture analysis for BDSS and GDSS. The curves fitted to the experimental 
data was appropriately adjusted with the Charpy impact energy curves expressed by a hyperbolic 
tangent function, as shown in Figure 7. The DBTT-FAT Ts that indicated 50:50 ductile behavior from the 
fractured surface of the specimens were 0.1 °C and −22.4 °C for BDSS and GDSS, respectively (Table 4). 
The DBTT value for BDSS, obtained from the FATT method, was almost 32 °C higher than that of 
the ETT method. Furthermore, the difference in the DBTT-FATT value between BDSS and GDSS was
greater than in the other ITT values, which included the DBTT-ETT. Some researchers have also found differences in the DBTT-ETT and DBTT-FATT values [69–71], suggesting that they could occur when the shape of the transition curve was unusual with an ambiguous transition point or at significantly low USE value [71]. By using a symmetric hyperbolic tangent function, as used for the Charpy impact energy fitting curves, the slope (C) of the transition region of BDSS had a larger standard error and a lower coefficient of determination, R-squared than that of GDSS.

Figure 10. The average fraction of ductile fracture obtained from fracture surfaces of BDSS and GDSS at various temperatures.

Microstructure and chemical composition of typical inclusions in each BDSS and GDSS are shown in Figure 11. The inclusions in dimples are clearly visible after Charpy impact test. The shape of inclusion was certainly different in both SDSSs. In the dimple of BDSS, the inclusions were angulated (Cr, Mn)-based oxides (Figure 11A,C), whereas relatively spherical shaped (Gd, Cr, Si)-based oxides were found in the dimple of GDSS (Figure 11B,D). During casting process, Gd acted as a flux and hence formed floating slags with large inclusions [10]. Gd also decreased the size of inclusions and changed the inclusions to be more spherical in shape, thus further improved the Charpy toughness of SDSS.
4. Conclusions

In this study, the effect of Gd addition on the microstructures and Charpy impact properties of super duplex stainless steels was investigated. A890 grade 7A SDSSs with and without Gd were successfully fabricated by an air-casting method. Reduction of the grain size and inclusion of SDSS were achieved by Gd addition because of a decrease in the nuclei formation’s activation energy and high reactivity of Gd with O. The addition of Gd to SDSS also resulted in the transformation of the non-metallic inclusions, and a reduction of their average fraction of inclusions. These combined effects improved the impact toughness and ductility of SDSS even though the relative number of inclusions increased and the average distance between them decreased. Compared with BDSS, GDSS demonstrated a higher absorbed energy, including LSE and USE, as well as a lower impact transition temperature (27J-ITT, 68J-ITT, and DBTT). Specifically, the DBTT calculated by FATT significantly decreased by more than 20°C upon the addition of Gd. This approach to improve microstructure and mechanical property of SDSS was effectively performed. Therefore, the modified SDSS by the addition of Gd have a great potential to become a structural alloy for applications under some extreme conditions.

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