Surface- and volume-based investigation on influences of different Varestraint testing parameters and chemical compositions on solidification cracking in LTT filler metals

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Received: 17 October 2019 / Accepted: 16 March 2020 / Published online: 27 March 2020
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Abstract
The subject of this study is how, and to what extent, Varestraint/Transvarestraint test results are influenced by both testing parameters and characteristics of evaluation methods. Several different high-alloyed martensitic LTT (low transformation temperature) filler materials, CrNi and CrMn type, were selected for examination due to their rather distinctive solidification cracking behaviour, which aroused interest after previous studies. First, the effects of different process parameter sets on the solidification cracking response were measured using standard approaches. Subsequently, microfocus X-ray computer tomography (μCT) scans were performed on the specimens. The results consistently show sub-surface cracking to significant yet varying extents. Different primary solidification types were found using wavelength dispersive X-ray (WDX) analysis conducted on filler metals with varying Cr/Ni equivalent ratios. This aspect is regarded as the main difference between the CrNi- and CrMn-type materials in matters of cracking characteristics. Results show that when it comes to testing of modern high-performance alloys, one set of standard Varestraint testing parameters might not be equally suitable for all materials. Also, to properly accommodate different solidification types, sub-surface cracking has to be taken into account.

Keywords Solidification cracking · Varestraint testing · LTT filler metal · Microfocus X-ray computer tomography (μCT)

1 LTT filler metals

The use of low transformation temperature filler metals is an innovative way to reduce tensile residual stresses that commonly occur when welding high-strength steels. Through addition of alloying elements, like nickel and manganese, the martensite start (Ms) temperature of these materials is considerably lower than for conventional filler metals. The increase in volume caused by the phase transition from austenite to martensite is hindered by the cold base material surrounding the weld, which leads to compressive stresses. This is counteracting the shrinkage-induced tensile stresses. The lower the Ms, the more pronounced this effect becomes. Tensile residual stresses can be reduced or even turned into compressive stresses.

The subject of low transformation temperature (LTT) has been intensively studied [1, 2]. In particular, the proof that compressive residual stresses are formed [3, 4], the investigation of the mechanisms of stress formation [5, 6], and the effect on the fatigue strength [7, 8] have been the subject of many research projects. Recent publications also deal with extended topics such as microstructure and the associated mechanical properties [9–13], the behaviour during multilayer welding [14–18], and the application of LTT in beam welding [19].

However, the chemical composition of the most commonly used LTT approach, with Cr and Ni as the main alloying elements, still involves the risk of solidification cracking. Only a few investigations are addressing this topic [20–22]. These show that the tendency for cracking is very sensitive to the chemical composition. Above all, the carbon and nickel contents determine the extent of crack formation.
The influence of the chemical composition of different LTT fillers on the tendency to solidification cracking is one of the main objectives of the present study. The different filler materials used in this regard stem from previous investigations aiming to evaluate the influence of the chemical composition on the $M_s$ temperature [21]. Their compositions are listed in Table 2 and shall be discussed in more detail later on.

As an example of the three alloy types examined in [21], Figs. 1, 2, and 3 show the cross sections produced by optical microscopy for multilayer welds. Lichtenegger and Bloech I etchant was chosen to contrast the weld microstructures. This etchant is well suited to reveal segregation in high-alloy welds. During manufacturing of the multilayer welds microcracks already appeared in individual cases (Figs. 1 and 3).

Alloys A and B, shown in Figs. 1 and 2, respectively, are both Cr and Ni based; their weld microstructures show similar characteristics. The structure is martensitic, which is etched blue and brown, and the micrographs give no indication of retained austenite. The prior austenite grain boundaries are partially visible. Some precipitations are present, independent of the varying carbon contents of the alloys. In the top layer of alloy A, a microcrack appeared along the boundary of two former austenitic cells. This location indicates a solidification crack. No further cracks were found in the cross sections.

Alloy G is Cr and Mn based, following a different LTT approach. Its weld microstructure is martensitic, which is etched brown as indicated in Fig. 3. Microsegregation is indicated by light and dark etched areas. Their appearance suggests a cellular structure, which is typical for primary austenitic solidifying welds. A microcrack was found in the weld root.

2 Solidification cracking and suitable testing methods

Solidification cracking is a defect connected to the solidification process during fusion welding of metals, caused by a combination of thermomechanical and metallurgical aspects. Certain alloying elements form low-melting phases in which cracks tend to initiate under deformation within a critical temperature range. In consequence, the solidification cracking susceptibility of a material can be described by its so-called brittleness-temperature-range (BTR) [24]. Cracking can only occur if strains within the critical temperature range exceed a certain threshold. The BTR therefore poses a solely material-specific parameter.

Under certain conditions, residual liquid contents flowing into previously formed cracks during further cooling can result in crack healing [25, 26]. For example, alloys with a composition close to the eutectic are known for low solidification cracking susceptibilities because of their high potential for crack healing [27].

The connection between primary solidification mode and solidification cracking susceptibility has been the subject of numerous studies. Primary austenitic solidification is known to have a negative influence on a material’s cracking susceptibility, and certain constituents of weld-ferrite are known to have a positive effect in this regard [28–30]. Compared with ferritic materials, slower diffusion and lower solubility for alloying elements in austenite promote segregation [26]. Furthermore, austenitic materials feature higher thermal expansion and coincidentally lower thermal conductivity, which also supports solidification cracking [31].

For solidification cracking testing, among other methods, the modified Varestraint/Transvarestraint test (MVT), developed at the Bundesanstalt für Materialforschung und -prüfung (BAM) [32], can be used. In this test, cuboid blocks are melted at the surface using a gas tungsten arc welding (GTAW) torch moving along the specimen. During this welding process, the specimen is bent over dies with different radii, length- or crosswise (Fig. 4). Through deformation to specific levels at specific speeds and within the solidification temperature range of the material, the general requirements for solidification cracking are met. Standard test evaluation relies on quantification of surface cracks, which can then be translated into an assessment of solidification cracking susceptibility of the material.

Experience from recent testing shows that some materials, particularly modern high-performance alloys such as LTT, also exhibit cracking in the bulk [20]. In those cases, the standard MVT procedure, which relies on surface cracks, might be insufficient for assessing the cracking susceptibility. The current study therefore aims to extend the evaluation basis to the volume of the specimen, allowing for a more reliable assessment of cracking susceptibility.
3 Influence of MVT testing parameters on the solidification cracking susceptibility

As inherent to the functional principles, applied strain is the key factor influencing the extent of cracking detected in Varestraint testing. Other main influencing parameters are heat input, travel speed, specimen thickness, and stroke rate (bending speed) [26, 33].

Numerous studies regarding the direct influence of testing parameters on the results have been conducted in the past, often with diverse results. While the general agreement is that higher strains increase the cracking response [24, 26, 33], contradicting opinions on the influence and general importance of other process parameters can be found in the literature.

Specimen thickness is usually not varied within a series of experiments, but regarded to be a significant influence on test results [34]. This aspect becomes even more complex when considering that nominal applied strain often deviates significantly from the actual local deformations in the vicinity of the weld pool, which pose the critical factor but at the same time are highly inhomogeneous [35].
Travel speed has been reported to be the most complex [34] or even critical [36] parameter, said to increase [37] or to decrease [38] cracking under otherwise equal conditions.

Some studies consider stroke rate to be of minor importance [36], while others regard it to be even more critical than travel speed [39]. It is furthermore expected to influence crack lengths due to crack extension during bending [40], and to delimit the amount of strain taking effect during a confined solidification interval [25]. In MVT-specific investigations, stroke rate was even found to have a linear correlation to cracking response, making it an important factor for quantifying cracking. Low stroke rates also appeared to favour centreline cracking, while increased stroke rates caused expansion of the crack afflicted area towards the edges of the weld [31].

In summary, Varestraint testing parameters are generally accepted to have a significant influence on test results, but still there is no general agreement on test specifications. Test results and parameters cannot be compared directly, and apparent contradictions cannot be discussed independently. This problem is further illustrated in Table 1, which is a review of technical standards, recent publications, and selected particularly reputable work, showing how much common Varestraint testing parameters diverge between different testing facilities.

The present study therefore aims to evaluate influences of different Varestraint testing parameters on solidification cracking in LTT alloys of different compositions. As sub-surface cracking has been observed in the past, volume-based analysis methods are to be employed in this regard.

4 Materials

Several Cr/Ni- and Cr/Mn-based LTT alloys were used in this study (see Table 2). Alloys A and B contain more Cr than Ni. In contrast, alloys C-F exhibit a higher Ni content compared with Cr. Alloys G and H are Cr/Mn alloyed, with Cr being the main alloying element. Mn shows a lower austenite stabilizing effect compared with Ni; therefore, these wires contain more Cr. Cr alone is ferrite stabilizing, but in combination with Mn or Ni, it causes a reduction of the M_s temperature. Variants with higher carbon contents have shown to be particularly susceptible to solidification cracking [20–22] and were therefore used in this study.

The Cr, Ni, Mn, and C contents were adjusted gradually in order to systematically vary the Cr_eq/Ni_eq ratios, as shown in Table 2. Other accompanying elements were not varied. The M_s temperatures have a range of 164 to 297 °C. All fillers were produced as metal-cored wires with a diameter of 1.6 mm.

5 Experimental

5.1 Standard MVT procedure

In order to apply MVT testing to filler metals, specimens must be made from deposited weld metal. Blank specimens were manufactured from S235 mild steel, featuring a rounded centre groove (5 mm deep, 20 mm wide, 5 mm corner radius; see Fig. 5). The groove was then filled with weld metal using gas metal arc welding. This was done in multiple passes in order to avoid any results being corrupted by the properties of the specimen base material. After welding, the specimens were machined and ground to match standard MVT dimensions (100 × 40 × 10 mm; see Fig. 5). During the following actual testing, a moving GTAW torch was melting only the deposited weld metal in the top centre of the specimen.

An investigation on MVT parameter sets outside of the standard range was carried out on alloys A and H, with a focus on stroke rate and energy input. This was expected to characterize the materials in more detail, therefore helping to find suitable testing parameters for future experiments on similar materials.

Rather, high stroke rates for Varestraint testing are the standard for many testing facilities around the world (see Table 1).
However, high stroke rates have been reported to suppress potential back-filling of cracks, making Varestraint testing less suitable for assessment of materials which might exhibit crack healing \[52, 53\]. Testing at different strain rates was therefore carried out in search for further insight into the crack healing potentials of the examined materials. In order to provide for comparability to previous studies at BAM, the current study uses a default stroke rate of 2 mm/s. Additional experiments were set at stroke rates of 0.2 mm/s and 6 mm/s, which is relating to the minimum and maximum machine settings.

Table 3 shows the range of parameters covered by the experiments. For the main set of experiments, 46 specimens made from alloy A and alloy H filler metals each (92 in total) were processed in 10 Varestraint and 10 Transvarestraint tests each, with 2–3 specimens per parameter set. From all other alloys, 2 specimens each were tested at 7.5 kJ, 2 mm/s stroke rate, and 4% strain. After testing, following the standard MVT analysis procedure, the specimens were examined by capturing the surface cracks that are visible at × 25 magnification through an optical microscope.

### Table 1: Common Varestraint test parameter sets found in the literature

| Source | Current (A) | Voltage (V) | Travel speed (cm/min) | Energy input (kJ/cm) | Stroke rate (mm/s) | Strain (%) | Magnification (optical microscope) | Specimen thickness (mm) |
|--------|-------------|-------------|-----------------------|----------------------|-------------------|------------|-----------------------------------|------------------------|
| ISO TR 17641-3 [41] | 85 | 12.5 | 18 | 3.54 | >1.8 | n/a | 25 | 10 |
| ISO TR 17641-3 [41] | 220 | 13.5 | 11 | 16.20 | >1.8 | n/a | 25 | 10 |
| ANSI AWS B4.0:2016 [42] | n/a | n/a | n/a | n/a | n/a | 0–4 | 40–80 | 3–10 |
| BAM Standard no. 1 | 190 | 12 | 18 | 7.60 | 2 | 0–4 | 25 | 10 |
| BAM Standard no. 2 | 210 | 12.7 | 10.8 | 14.82 | 2 | 0–4 | 25 | 10 |
| Zhu et al. [43] | 150 | 17 | 10 | 15.30 | n/a | 0–6 | n/a | 6 |
| Statharas et al. [44] | 225 | 12 | 21 | 7.71 | n/a | 2–11 | n/a | 24 |
| Singh et al. [45] | 180 | 10 | 13 | 0.80 | 1 | 5 | n/a | 5 |
| Abe et al. [46] | 200 | 14.3 | 16 | 10.73 | 150 | 1.96–5.88 | n/a | 10 |
| Singh et al. [47] | 70 | n/a | 6 | n/a | 10 | 0.8–8.2 | n/a | 3.3 |
| Lippold [26] | 160–190 | n/a | 10.2–15 | n/a | 152–254 | 0.5–7 | 20–50 | 6.35 |
| Srinivasan et al. [48] | 90 | 11.5 | 25.2 | 2.46 | Within 0.5–4 | 0.015 s | 60 | 3 |
| Andersson et al. [36] | 70–90 | n/a | 6–18 | n/a | 10–250 | 2.7–4 | n/a | 3.2 |
| Yushchenko et al. [49] | 90 | 9.7 | 4.8 | 10.91 | n/a | 0.2–2 | 50 | 3.5 |
| Lundin et al. [34] | 190 | n/a | 12.7 | n/a | Within 0.1 s | 0–4 | 40–80 | 7.925 |
| Arata et al. [50] | 250 | 17 | 10 | 25.50 | n/a | 0.2–4 | 40 | 12 |
| Senda et al. [51] | 250 | 18 | 15 | 18.00 | 170–250 | 0.15–1.5 | 42–100 | 6–9 |

### Table 2: Chemical composition of the LTT alloys used in this study, determined by spectral analysis (wt%)

| Type key | C  | Cr  | Ni  | Mn  | Fe  | Cr<sub>eq</sub> | Ni<sub>eq</sub> | M<sub>l</sub> in °C |
|----------|----|-----|-----|-----|-----|--------------|---------------|---------------|
| CrNi A   | 0.05 | 7.48 | 6.05 | 0.47 | bal. | 0.98 | bal. | 297 |
| CrNi B   | 0.07 | 7.41 | 6.08 | 0.49 | bal. | 0.90 | bal. | 275 |
| CrNi C   | 0.07 | 6.30 | 7.90 | 0.56 | bal. | 0.61 | bal. | 268<sup>b</sup> |
| CrNi D   | 0.09 | 7.10 | 8.20 | 0.60 | bal. | 0.64 | bal. | 238<sup>b</sup> |
| CrNi E   | 0.10 | 6.80 | 7.70 | 0.60 | bal. | 0.63 | bal. | 247<sup>b</sup> |
| CrNi F   | 0.11 | 6.50 | 7.90 | 0.59 | bal. | 0.56 | bal. | 245<sup>b</sup> |
| CrMn G   | 0.07 | 11.90 | 0.02 | 5.40 | bal. | 2.47 | bal. | 164 |
| CrMn H   | 0.03 | 11.40 | 0.02 | 5.00 | bal. | 3.28 | bal. | 176 |

<sup>a</sup> Calculated as follows: Cr<sub>eq</sub> = Cr, Ni<sub>eq</sub> = Ni + 30C + 0.5 Mn
<sup>b</sup> Calculated after [23]

**Fig. 5** MVT specimens before deposition of weld metal (left) and after finishing, ready to test (right)
of cracking susceptibility, volume information should also be considered. Therefore, microfocus X-ray computer tomography (μCT) was applied to crack afflicted areas of selected specimens at Karlsruhe Institute of Technology (KIT) using the high-resolution cone-beam CT system type Yxlon Y.CT Precision with fine focus twin head FXE 225.99. These scans were performed at a high voltage of 190 kV and 0.06 mA. This allows characteristics of the complete 3D crack network to be extracted.

The first step during μCT analysis consisted of scanning each specimen as a whole, in order to locate the area of interest for exact quantification of volume cracking. This provides a good overview of crack afflicted areas, which after localization can be physically extracted from the large specimens. The now considerably reduced absorption thickness (about 10 mm) leads to significantly improved image qualities. As the smaller specimens can also be positioned closer to the X-ray source, spatial resolution of the reconstructed 3D images is improved due to the cone-beam geometry (uncut specimens ~ 25 μm voxel size, cut specimens ~ 7 μm voxel size). At this point, the obtained image quality is still not sufficient for reliable and fast segmentation of the cracks using commercially available software. In this regard, a two-step segmentation strategy was implemented. In the first step, an adaptive thresholding algorithm (described in [54]) was applied. Here, the 3D image is partitioned into small cubes with defined edge length. Within the cuboids, a mean local grey value is calculated. If the grey value of a voxel is below a certain threshold of the mean grey value (e.g. 10%), this voxel is detected as a crack. The adaptive thresholding algorithm allows handling of the large grey value fluctuations within the reconstructed 3D specimen volume. In the second step, the segmented voxels from the first step are used as the so called seeds for a region growing algorithm [55]. This segmentation strategy examines neighbouring voxels of the initial “seeds” and determines by means of a second threshold whether they should be added to the crack region or not. Using the region growing algorithm allows the growing or shrinking of the detected cracks to their final size. Subsequently, the segmented binary images are analyzed statistically and quantitatively. Therefore, the 3D crack network is described in regard to e.g. crack volume, cracking surface, crack length over depth, crack orientation, or degree of branching.

The μCT analysis procedure takes about 6 h in total per specimen, including specimen preparation (cutting), 2 h of scan time, and data evaluation.

### 6 Results

#### 6.1 Standard MVT analysis

A variation of total strain was performed for alloys A (CrNi type) and H (CrMn type). For both alloys, strains of 1% did not result in any cracking. Further increasing of the strain to 4% caused cracks to form, to different extents. Therefore, only experiments set at 4% strain will be considered below.

In the same way, no cracks could be detected for stroke rates of 0.2 mm/s, regardless of material composition or other process parameters. Thus, only experiments at stroke rates of 2 mm/s or above will be considered from here on.

All observed cracks were located within the weld and identified as solidification cracks. Their exact location varies slightly between Varestraint and Transvarestraint tests, with the former producing a more spread-out crack pattern, while the latter favoured cracks closer to the weld centreline. The degree of surface cracking is herein after assessed by means of total crack length (TCL), which was measured according to the MVT standard analysis procedure.

Figure 6 shows the total crack lengths for all alloys at 7.5 kJ/cm energy input, 2 mm/s stroke rate, and 4% strain. Among all alloys, the CrMn types (G and H) show the lowest extent of cracking. CrNi-type alloys B and C exhibit around 4 mm TCL; all other alloys are above 9.88 mm TCL.

The influence of different MVT testing parameter sets on the results was investigated for alloys A and H; results are shown in Fig. 7. Contradicting trends were found concerning the effects of energy input and stroke rate.

Alloy A shows a considerable degree of cracking for lower energy input in Varestraint tests at 2 mm/s stroke rate, while at high energy input, almost no cracking occurs. Increasing the stroke rate to 6 mm/s practically levels the influence of energy input. Transvarestraint testing produces very uniform results for all parameter sets. Just as in the Varestraint tests, lower energy input causes higher cracking response, although the difference is far more subtle. In general, however, neither energy input nor stroke rate seems to have significant influence on the Transvarestraint test results, as the TCL remains at a very similar level.

The Varestraint results for 2 mm/s stroke rate and 7.5 kJ/cm energy input seemed out of line with the general trends. In order to validate the results in question, an additional experiment was carried out later on as a follow-up to the original experimental plan. Three specimens were tested at 10 kJ/cm
energy input (2.4 mm/s travel speed, 200 A), 4% strain, and 2 mm/s stroke rate. The result falls right in between the other data points.

Alloy H shows a much more consistent response to different testing parameters, and no qualitative difference between Varestraint and Transvarestraint modes. Higher energy input generally causes a higher degree of cracking, regardless of stroke rate. Higher stroke rates cause a significant but nonlinear increase in cracking. This seems to reduce the influence of energy input, as the TCL at different energy inputs converges for higher stroke rates.

Metallographic cross sections for alloys A and H are shown in Fig. 8. It can be seen that alloy A is exhibiting a considerable amount of cracking below the surface, while cracks in alloy H generally tend to propagate to the specimen surface, displaying an amount of visible cracking that is more in proportion to crack volume below the surface.

### 6.2 μCT analysis approach

In order to address the different behaviours concerning crack propagation into the bulk (Fig. 8), further investigation in this regard was conducted on all alloys using μCT analysis. Unfortunately, due to inadequate specimen preparation, no results could yet be obtained for alloy D.

Figure 9a shows TCL at different depths for alloys C and E, calculated from μCT scans. It can be seen that the total crack length at the surface (0 mm depth) is very similar, around 2.5 mm for both materials. However, cracks in alloy E reach much deeper into the specimen volume than in alloy C, around 3.0 mm versus 2.5 mm, respectively.

Even more significant differences become apparent when comparing alloy B and C weld metals (Fig. 9b), featuring 2.6 mm and 0.75 mm total crack length at the surface,
respectively. Looking into the volume, cracks in alloy B reach deeper than in alloy C (2.8 mm versus 2.5 mm), even though surface-based evaluation clearly presents alloy B to be less prone to cracking.

Investigations on CrMn-type alloys turned out to be more challenging, as both alloys G and H exhibit very little cracking. In addition, the morphology of crack networks is considerably finer than for CrNi-type alloys, making them increasingly difficult to detect. For the case of alloy H, no cracks could be found in the μCT investigations, while the standard analysis clearly showed surface cracking, albeit at a very low level (see Fig. 11).

7 Discussion

Higher energy inputs were expected to have a negative effect on the degree of solidification cracking. In the case of alloy H, this is reflected by the MVT standard analysis results. Alloy A exhibits an entirely different behaviour, with crack length decreasing at increased energy input. Due to the different nature of the two alloy types (CrMn and CrNi, respectively), further examination of the solidification characteristics and the specimen volumes was expected to generate more insight.

In accordance with analytic calculations after Hull [56] and Hammar and Svensson [57], the WDX analysis of alloy A and alloy H filler metals indicates different primary solidification modes. The microstructure of alloy A (Cr(eq)/Ni(eq) = 0.98) is marked by concordant distribution of Cr and Ni (Fig. 10a), which is typical for austenitic primary solidification. The alloy H weld metal (Cr(eq)/Ni(eq) = 3.28) on the other hand exhibits discordant distribution of Cr and Mn (Fig. 10b), indicating ferritic primary solidification [28, 58].

Crack volume determined in μCT analysis on MVT specimens is shown in Fig. 11 for all alloys as a function of their Cr(eq)/Ni(eq) ratios (blue line, alloys from left to right: F, C, E, B, A, G, H). It can be seen that with increasing Cr(eq)/Ni(eq) ratio, the crack volume decreases. For Cr(eq)/Ni(eq) ratios of 3 and above, no cracks are found, while a peak level of around 3.3 mm³ is reached for Cr(eq)/Ni(eq) ratios between 0.55 and 0.62. The reason for this is expected to be the austenitic primary solidification mode prevailing for Cr(eq)/Ni(eq) ratios < 1.5. This level has been reported to be a threshold above which a minimum cracking susceptibility is to be expected in the case of stainless steels [28, 59].

Figure 11 furthermore shows a comparison between total crack length found in MVT standard analysis and crack volume found in μCT analysis, again referenced to the corresponding Cr(eq)/Ni(eq) ratios (red line, alloys from left to right: F, C, E, D, B, A, G, H). While the results from both analysis methods match the presumed coherence between cracking.

Fig. 8 Solidification cracks in alloy A (a) and H (b) from Transvarestraint specimens tested at 4% strain, 15 kJ/cm energy input, and 6 mm/s stroke rate; cross sections were taken from the crack afflicted areas in the specimen centre, normal to the welding path.

Fig. 9 Total crack length as a function of depth, evaluated using μCT analysis for alloy C and E weld metals (a), as well as alloys B and C (b).
response and Cr\textsubscript{eq}/Ni\textsubscript{eq} ratio, there is a more consistent trend for the μCT results. This is attributed to the significant amount of cracking in the specimen volume which was observed for CrNi-type alloys (see Fig. 8) but is not considered in the standard analysis process. Therefore, assessment of cracking susceptibility based on 3D crack networks brings a considerable added benefit over the conventional, surface-based analysis procedures.

Regarding the variation of testing parameters which was carried out for alloys A and H, different factors were found to trigger cracking. For the case of alloy A, cracking is substantially favoured by low energy inputs. μCT analysis is able to back up this finding. This behaviour was attributed to conflicting influences of weld pool geometry and solidification characteristics for varying energy inputs or welding speeds [60], yet additional investigations are required to fully explain these results. Cracking in alloy H was found to be triggered by higher stroke rates. The reason for this is believed to be the alloy’s considerable ability to back-fill existing cracks, given that there is enough time. This also explains why regardless of any other parameters, no cracking was observed for the extremely low stroke rate of 0.2 mm/s.

8 Summary and conclusions

Different CrNi- and CrMn-type LTT filler metals were examined regarding their solidification cracking susceptibility. In addition to standard Varestraint analysis methods, a μCT approach was used to quantify cracking in the specimen volume. The following conclusions can be drawn:

- Most of the evaluated alloys showed significant, yet varying extents of cracking in the specimen volume. Surface-based analysis methods therefore entail inaccurate representation of cracking susceptibilities. Volume-based examination should be used for better quantification in these cases.
- The μCT analysis approach discussed in this study was proven to be a suitable method for more accurate, volume-based specimen evaluation.
- With increasing Cr\textsubscript{eq}/Ni\textsubscript{eq} ratios, the 3D crack networks become increasingly finer in morphology. Because of its resolution limit, the μCT analysis in its current state of development works best on those LTT alloys with Cr\textsubscript{eq}/Ni\textsubscript{eq} ratios of 2.5 or below.
- The evaluated alloys with a Cr\textsubscript{eq}/Ni\textsubscript{eq} ratio of 1.5 or lower show the highest cracking susceptibility, which is in good agreement with the literature on common stainless steels.
- The CrNi-type alloy A is almost insensitive to differences in stroke rate but decreasing energy input favours cracking. Additional specimens need to be tested in order to back up and explain this finding in more detail.
- Cracking of the CrMn-type alloy H is predominantly triggered by increasing stroke rates, which is attributed to a higher potential for crack healing.
• Common Varestraint test parameters are highly inconsistent, yet different material characteristics might call for more specific test conditions in certain cases. As some of the investigated materials seem to bear considerable potential for crack healing, an adjusted variation of stroke rates shall be used in future investigations on this matter.

**Funding information** Open Access funding provided by Projekt DEAL. This research received financial support from the German Research Foundation (DFG) in the projects GI376/8-1, KA1807/9-1, and KR3917/2-2.

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