



HYDROSTATIC TESTING OF WELDED STAINLESS STEEL FABRICATIONS



Application Data Sheet – Stainless Steel

Hydrostatic Testing of Welded Stainless Steel Fabrications

Selection and Treatment of Water and Conduct of the Test, to Maintain the integrity of the System

1.0 Introduction

Welded fabrications are frequently tested hydrotested for pressure tightness. Unless properly conducted, this procedure can induce problem initiating corrosion of the fabrication by subjecting it to critical conditions outside the initial design constraints. Where the vessel is to handle ingested, contamination must be avoided.

Damage may be caused by the induction of latent corrosion problems during and after hydrotesting. The damage may not become apparent months later, and may very difficult and costly to repair.

However, if simple precautions are taken, hydrotesting can be conducted without risk to the equipment. The precautions mainly relate treatment of the water used, and how the equipment is treated after hydrotesting. And hydrotesting even offers an opportunity to improve fabrication during testing.

This note gives general advice on how to conduct hydrotesting without damaging the equipment. However, the subject is complex, and the tester to ensure that the practices used will not cause problems. There are no published standards or codes of conduct reporting industry codes for stainless steels, although several give general advice (see boxes below and on page 8). Please consult Austral Wright Metals for further information.

2.0 Corrosion of Stainless Steels

Corrosion failures of fabricated stainless steel systems soon after hydrotesting may result from one or more of three causes:

- Pitting & crevice corrosion (PCC)
- Stress corrosion cracking (SCC)
- Microbially induced corrosion (MIC)

API 570: Piping & Inspection Code, American Petroleum Institute.

Piping fabricated of or having components of 300 series stainless steel should be hydrotested with a solution made up of potable water (see condensate). After testing is completed, the piping should be thoroughly drained (all high-point vents should be opened during draining), air dried. If potable water is not available or if immediate draining and drying is not possible, water having a very low chloride level, higher pH (addition may be considered to reduce the risk of pitting and microbially induced corrosion).

Note: Potable = 'suitable for drinking'. Potable water in this context follows US practice, with 250 ppm maximum chloride, sanitized with chlorine.

Pitting and crevice corrosion can result from attack by the water used for hydrotesting. The equipment is designed for particular conditions. Surface finish are selected from knowledge of the proposed temperature, solute concentrations, pH etc. If the water conditions used for test

corrosion may occur. Unfortunately, corrosion by these mechanisms is localised, and proceeds in isolation from the bulk solution conditions proceed in conditions which are otherwise not aggressive. Failure may occur by local penetration 3 – 12 months after initiation of the corrosive

The most aggressive ion in water is usually chloride, and accepted limits of this ion to avoid pitting & crevice corrosion are 200ppm for grade 316(L) and 4,000 for duplex 2205, see figure 1. These values are not conservative. Note that some authorities limit the concentration of chloride

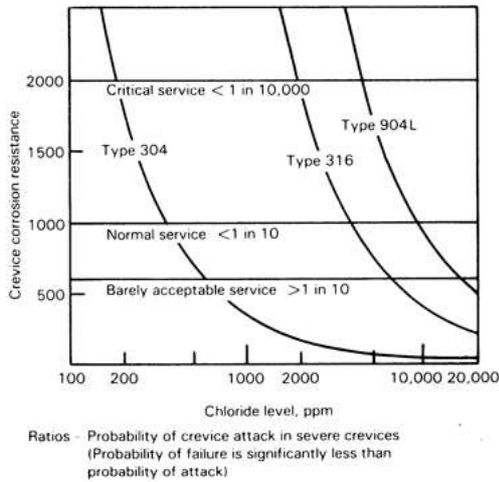


Figure 1: Probability of crevice corrosion attack as a function of chloride level. The line for the super austenitic 904L also represents the duplex grade 2205

Time and temperature are also important in the development of pitting & crevice corrosion, and water with higher chloride content may be used where temperature are limited.

Apart from the initial quality of the water, its aggressiveness can be increased by increasing temperature, by oxygen starvation under particle attack, and by concentration of chlorides in pools of water left in the system after testing. See section 5 for precautions to be taken after hydrostatic testing.

Stress corrosion cracking is a failure mechanism which can cause failure very rapidly. Stainless steels are not especially susceptible to stress corrosion cracking. Failure may be encountered at high chloride levels and temperatures in austenitic grades such as 304 and 316. The duplex grades (e.g. 2205) are effectively immune to failure in waters.

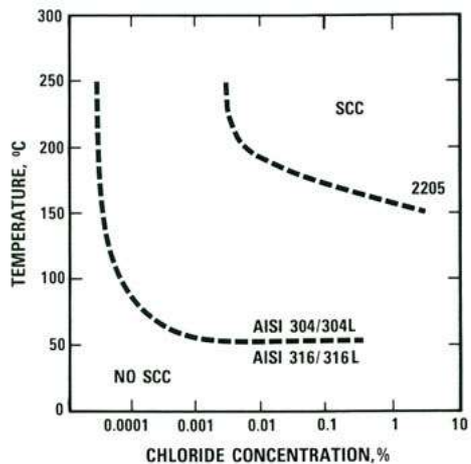
The austenitic grades are also effectively immune to stress corrosion cracking in waters at temperatures below about 50°C (figure 2). Since hydrostatic testing is often run at ambient temperature, the problem is unlikely to arise. However, equipment left in strong sunlight for a long period could get to this temperature. Care should be taken to avoid excessive temperatures.

Figure 2: Stress corrosion cracking of austenitic and duplex grades as a function of chloride concentration and temperature [3].

Note:

Chloride Concentration	
%	ppm
0.0001	1
0.001	10
0.01	100
0.1	1,000
1	10,000
10	100,000

Seawater contains ~19,000 ppm (2%) chloride



Microbiological Corrosion (MIC) is a mechanism which is increasingly being recognised as contributing to corrosion of metals [4]. Penetration rates have been reported for 304L, and in four months for 316L [5]. Micro-organisms, primarily bacteria, can alter the micro-environment to which they are exposed, concentrating aggressive species, and by restricting the access of oxygen to the surface.

Sulphate reducing bacteria (anaerobic) metabolise sulphate ions to sulphide ions, while sulphate oxidising bacteria (anaerobic) can metabolise organic acids. Both are highly aggressive to stainless steel.

The most common MIC problems with stainless steels are caused by iron utilising bacteria, which concentrate chloride as iron and manganese. These concentrations can easily far exceed the chloride levels mentioned above, and the resulting corrosion

and/or crevice corrosion: crevices, especially at gaskets, are one favoured site for this form of corrosion. Another common site is at the weld that properly treated welds are not generally preferential sites for corrosion, but it is probable that surface imperfections such as oxide or sludge *cetera* may be easy sites for bacteria to colonise.

MIC almost invariably occurs with raw (untreated) fresh water sources such as reservoirs, rivers, bores or wells, or fresh waters inadequately treated. Stagnant and low flow conditions promote attachment of bacteria to the metal surface. This is followed by colonisation and formation of discrete and frequently developing into a bulky deposit. The composition of the deposit is significantly different from and more corrosive than the bulk metal as [4]:

- large subsurface cavities or tunnels, frequently at or adjacent to welds
- as broad, open pits in crevices such as gasketed flanged joints
- as broad conical pits with terraced sides

MIC can be prevented by draining and drying equipment after hydrotesting. The bacteria require time to grow, and no significant damage should occur during the time of hydrotesting.

This supposes that drying is complete and thorough, with no possibility of pools being left in the system. It is more conservative to also treat the metal before use. This is discussed in more detail in section 4.

3.0 Water Quality

NHMRC/ARMCANZ *Australian Drinking Water Guidelines* (ADWG) give a guideline maximum value for chloride of 250 ppm, and most potable water should adhere to this limit. However, some supplies do not, especially where there is a significant proportion of bore water in the supply, and at times of drought may use alternative supplies. The Water Authority has information on the chloride level of the water supplied.

Since the 200 (304), 2,000 (316) and 4,000ppm (2205) chloride levels discussed above as being acceptable for avoiding pitting & crevice corrosion service, use of water somewhat above these levels should not cause problems at the shorter times used in hydrotesting. Moving from the 'service' line in figure 1 gives limiting chloride content of 400 ppm for 304, 4,000 ppm for 316, and 8,000 ppm for 2205. Obviously, this requires the equipment to be drained and dried after testing – see next section.

Other properties of potable water, such as pH, dissolved oxygen and hardness, are unlikely to cause corrosion problems for **stainless steel**.

4.0 Water Treatment

Where water is to be obtained from a **source other than a reticulated potable supply**, the chloride levels given above should still be adhered to. Water analysis labs will advise on how to take samples, and will analyse the samples for a hundred dollars or so. They may be found under "A" in the Telephone Directory.

Where it is essential to use water sources with chloride contents higher than these limits, and also for non potable supplies, the pH should be adjusted to say 7.5 – 10. Caustic soda (NaOH) can be used for this, and has a double benefit in that the hydroxide ion inhibits corrosion of **stainless steel**. Sulphuric acid may be added to low sulphate, high chloride waters, up to about 100 ppm.

Austral Wright Metals are able to predict the corrosivity of waters, using software developed by the Nickel Development Institute. Contact Austral Wright Metals for the software, or predictions of corrosivity are available to customers, requiring the input data as shown in Appendix 1 to produce the outputs.

Potable water supplies should be free of particulates, but other sources should be screened or settled and filtered. A filter passing particles of 10 microns is adequate.

Sanitisation or disinfection of the water is essential to remove the risk of MIC. It can be carried out by a variety of methods, some of which are more aggressive for other forms of corrosion. Potable water from properly conditioned town supplies do not require further sanitisation.

The most common sanitisation method is chlorination, and the most common method of introducing chlorine is with a hypochlorite salt, which is available as a liquid solution, most conveniently from swimming pool supply shops. The addition required is usually small, and online additions are difficult, so it is possible, to disinfect the water as a batch before introducing it to the system under test, adding the disinfectant last and ensuring good mixing. The quantity of sodium hypochlorite required for specific volumes of testing water. More polluted water supplies may need more hypochlorite. This should be checked with a normal colorimetric swimming pool chlorine test kit.

Free chlorine levels at entry to the system should be limited to 2ppm for grade 304, 5ppm for grade 316 [7]. The 2 ppm limit for residual free chlorine applies to copper alloys in the system.

Short term levels of free chlorine of up to 25 ppm can be tolerated by 316, provided there is thorough post cleaning treatment rinse, using water of the quality maxima already mentioned [7].

After hydrotesting it is prudent to measure the chlorine residual of the last water to be drained from the system. If this is below 0.2ppm, the system should be flushed with properly treated sanitised water without delay.

Other disinfectants which may be used include chlorine gas, chlorine dioxide, ozone and ultraviolet light. The former are subject to the same risks as residual disinfectants, while ozone and ultraviolet light pose no risks for stainless steel. Indeed, ozone may be beneficial for corrosion resistance.

Table 1: Quantities of sodium hypochlorite required for residual disinfectant in flushing & test water [6].

Volume of Test Water	Volume of 12.5% Sodium Hypochlorite to produce 2 ppm residual free Chlorine in Clean Water
Litres	Millilitres
10	0.2
50	1.0
100	2.0
1,000	20
10,000	200
50,000	1000

5.0 Precautions after Hydrotesting – Drying

Equipment left wet after testing is at risk because the chloride levels of pools of water will rise as the water evaporates: PCC may initiate at these levels. To prevent chloride concentration after hydrotesting, the equipment should be thoroughly dried and sealed if it is to be left for more than a short period of service. This can be achieved by draining the water out (properly designed equipment will allow for complete drainage), then blowing dry air through the system. If the humidity of the air is high, this process should be completed when the work is at its coolest (i.e. at night), to prevent condensation when the work is dried. Drying the equipment also stifles any microbiological infections, which will not progress on a dry surface.

6.0 Possible Beneficial Effects of Hydrotesting

ASTM A967 “Chemical passivation treatments for stainless steel parts” [8] gives a series of treatments for improving the corrosion resistance of stainless steel parts. Passivation is defined as the chemical treatment of a stainless steel with a mild oxidant, such as a nitric acid solution, for the purpose of removing surface foreign matter, but which is generally not effective in removal of heat tint or oxide scale on stainless steel.

By mixing a suitable oxidant with the hydrotesting water to passivate the surface, it should be possible to improve the overall corrosion resistance of stainless steel where a passivation treatment has not previously been applied to the surfaces. These passivation treatments require that the surfaces be free of oil, grease, rust, scale and other foreign matter.

The standard specifies five alternative treatments with nitric acid, but the most practical in this context is 20 to 45% nitric acid at 21 to 32°C followed by a clean water rinse.

Probably more practical for hydrotesting is one of the five citric acid based solutions, using 4 to 10% citric acid for a minimum of 20 minutes at 21 to 32°C. Care should be taken to evaluate the likely effect on the entire system: there may be carbon steel or copper alloys which would be unduly attacked. After testing, the system should be thoroughly rinsed, with water of adequate quality as detailed above. The acid effluent must be neutralised before discharge. It is good practice to dry the system within a day or two after testing.

While these treatments offer potential improvement to the corrosion resistance of the steel surfaces contacted, they are by no means essential and are outweighed by the costs of the chemicals, effluent neutralisation and rinsing.

7.0 Treatment of the Stainless Surface before Hydrotesting

In all the discussion in this note it has been assumed that all welds in contact with the hydrotesting water have been made or treated to remove welding oxide and adjacent chromium depleted zone from the surface. This is equivalent to external surfaces on the weld seams of Quality AS/NZS 1554.6:1994, “Welding stainless steels for structural purposes” [9]. The Grade II finish may be achieved by (a) acid pickling, (b) stainless steel finishing to a specified abrasive grit size, or (c) inert gas purging.

Where option (c), finishing, is specified, best corrosion performance is obtained with a surface roughness (Ra) of 0.5 micrometres or smoother. A worn belt of grit size 180#, although 220# is more certain.

Option (d) provided in AS/NZS 1554.6 of abrasive grit blasting is not recommended.

Where option (e) inert gas purging is used, AWS D18.1 “Specification for welding of austenitic stainless steel tube and pipe systems in situ” provides guidance and colour standards on the level of oxidation of the surface which is tolerable.

ASTM A380 “Standard practice for cleaning, descaling and passivation of stainless steel parts, equipment and systems” [11] provides detailed procedures for stainless steel surfaces for best corrosion resistance.

There is a reference to hydrotesting in the ASME Power Piping Code B31.1, relative to **stainless steel**. It appears in Appendix IV under Nonmar ASME B31.1 Power Piping Systems”, section IV-3.4, precautions for hydrotesting.

SUMMARY

- eliminate or at least minimise crevices in design & fabrication

slope horizontal lines and components to allow self draining, and provide adequate drainage at high and low points of the system

- use non wicking, chloride free gaskets at flanged connections
- specify full penetration welds, and monitor their achievement
- specify weld surface treatment to at least Quality B, Surface Finish II, as defined in AS/NZS 1554.6:1994. The finish may be achieved steel wire brushing, (c) buffing or finishing to a specified abrasive grit size (180# or smoother), or (e) inert gas purging.
- use the cleanest water available – demineralised, steam condensate, potable treated water if possible
- where the water does not come from at least an existing potable supply, analyse it to determine the corrosivity. Limit the chloride exposed, and adjust the chemistry (sulphate, pH) of the water. Also settle and/or filter it, and sanitise it to minimise the possibility of MIC.
- ensure sanitisation leaves a chlorine residual not exceeding 2 ppm for 304 or 5 ppm for 316 of chlorine at entry to the system
- prepare the water by batch treatment, adding the disinfectant last
- prepare the water by batch treatment before entry to the system
- regardless of water quality, drain, dry and inspect to assure dryness immediately following hydrotest, i.e. within three days
- If the residual free chloride is below 0.2 ppm as the testing water is drained from the system, or the water contains sludge or particles immediately with properly prepared, clean sanitised water

References

- [1] Nickel **stainless steels** for marine environments, natural waters and brines, Nickel Development Institute No 11 003, 1993.
- [2] RP0182-95 Standard recommended practice for initial conditioning of cooling water equipment, NACE, 1995.
- [3] A.J. Sedriks, Corrosion of **stainless steels**, 2nd Edn, Wiley, 1996.
- [4] G. Kobrin et al, Microbiologically influenced corrosion of **stainless steels** by water used for cooling and hydrostatic testing, Nickel Development Institute No 11 003, 1993.
- [5] G. Kobrin, Material Performance, vol 15, No 7, p40, July 1976.
- [6] DR 02014, Copper pipe and fittings – installation and commissioning, Standards Australia 2002.
- [7] **Stainless Steel** Advisory Service Information Sheet No.4.92, Issue 02, March 2001, British **Stainless Steel** Association, UK.
- [8] ASTM A967 Chemical passivation treatments for **stainless steel** parts.
- [9] AS/NZS 1554.6:1994 Welding **stainless steels** for structural purposes, Standards Association of Australia, 1994.
- [10] American Welding Society Standard AWS D18.1 Specification for welding of austenitic **stainless steel** tube and pipe systems in sanitary (hygienic) applications, 2002.
- [11] ASTM A380 Standard practice for cleaning and descaling **stainless steel** parts, equipment and systems.

Appendix 1

Nickel Development Institute Crevice Corrosion Model

a) Results screen

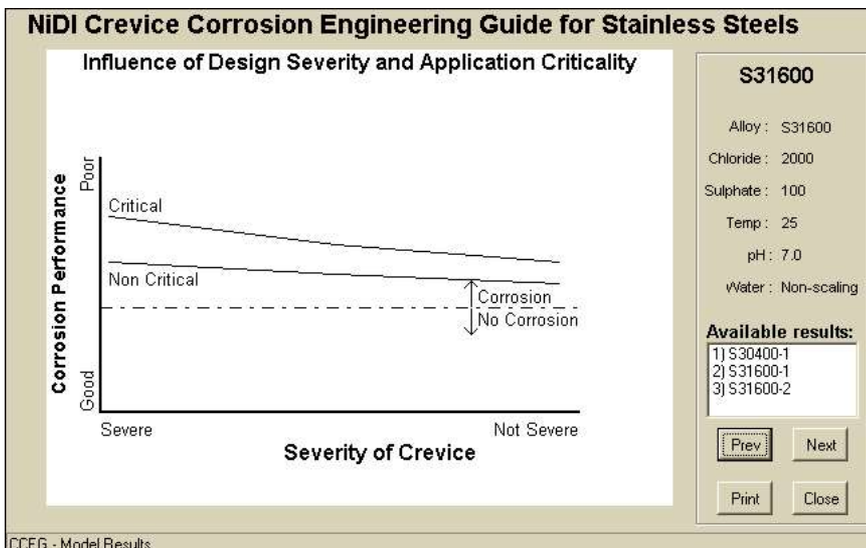
CCEG MODE 1 SUMMARY

The user has asked the CCEG for guidance on the corrosion of S31600 under the conditions specified below.

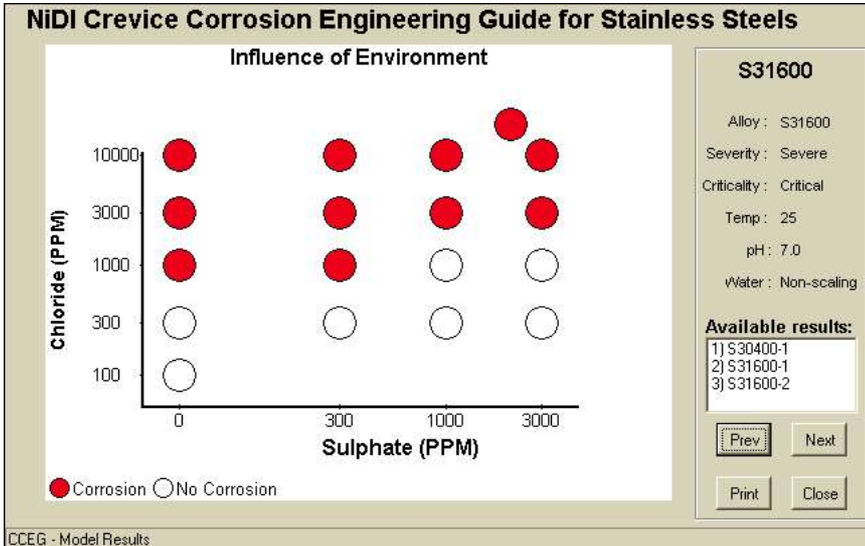
Under these conditions the CCEG predicts that corrosion will initiate in DAYS.

SERVICE APPLICATION		
Design Severity: Severe	Application Criticality: Critical	
WATER PARAMETERS		
Chloride (ppm): 2000	Sulphate (ppm): 100	
Hardness (as CaCO3 mg/L): 750	Alkalinity (as CaCO3 mg/L): 100	
TDS (mg/L): 3448	pH: 7	
Temp (C): 25	Scaling pH: 7.00	Water is N
Oxygen level (ppm): 7		
MATERIAL		
Form: Default (Wrought)	Fabrication: Welded	Condition:

b) Output Screen 1



c) Output Screen 2



Data for Crevice Corrosion Predictor

Austral Wright Metals Hydrotesting of Welded Fabrications

Data for Crevice Corrosion Predictor

Project: _____ Date: _____
 AWM Customer: _____ Phone: (0) _____
 Customer Contact Name: _____ Mobile: _____
 Email: _____ Fax: (0) _____

Tick boxes as appropriate

A: Stainless alloys used: 304 316 2205 Other (Specify) _____

B: Form: Wrought Cast

C: Surface: As received Ground Pickled Electropolished

D: Welded: Yes No

E: Welds treated by: Ground Pickled Both

F: Water Quality:
 pH: _____ Sulphate (ppm): _____
 Chloride (ppm): _____ Alkalinity (ppm): _____
 Total Hardness (ppm): _____ Conductivity (µS/cm): _____
 Total Dissolved Solids (ppm): _____

Form completed form to (02) 9627-6745, Attention Alex Gough.
 Austral Wright Metals supplies a comprehensive range of stainless steels, copper alloys and other performance metals for hydrotesting.
 Page 1 of 1

The technical advice and recommendations made in this Product Data Sheet should not be relied or acted upon without conducting your own including corrosion exposure tests where needed. Please consult current editions of standards for design properties. Austral Wright Metals with the information in this Product Data Sheet. Austral Wright Metals supplies a comprehensive range of stainless steels, copper alloys, nickel performance metals for challenging service conditions. Our engineers and metallurgists will be pleased to provide further data and applicatio

QUICK NAVIGATION

Home

Locations

Enquiry Form

Terms & Conditions

[Privacy Policy](#)

LOCATIONS

Sydney NSW
Melbourne VIC
Adelaide SA
Perth WA

SEARCH AWM WEBSITE

What are you looking for?

COPYRIGHT © AUSTRAL WRIGHT METALS :: AUSTRALIA 2000-2015 | AW DISTRIBUTION PTY LTD | ALL RIGHTS RESERVED | SITE BY I.T.B