Effectiveness of Four Disinfectants against Ebola Virus on Different Materials

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Abstract: The West Africa Ebola virus (EBOV) outbreak has highlighted the need for effective disinfectants capable of reducing viral load in a range of sample types, equipment and settings. Although chlorine-based products are widely used, they can also be damaging to equipment or apparatus that needs continuous use such as aircraft use for transportation of infected people. Two aircraft cleaning solutions were assessed alongside two common laboratory disinfectants in a contact kill assay with EBOV on two aircraft relevant materials representative of a porous and non-porous surface. A decimal log reduction of viral titre of 4 is required for a disinfectant to be deemed effective and two of the disinfectants fulfilled this criteria under the conditions tested. One product, Ardrox 6092, was found to perform similarly to sodium hypochlorite, but as it does not have the corrosive properties of sodium hypochlorite, it could be an alternative disinfectant solution to be used for decontamination of EBOV on sensitive apparatus.

Keywords: Ebola virus; disinfection; decontamination; inactivation; sodium hypochlorite; safety

1. Introduction

The worst outbreak of Ebola virus disease (EVD) started in Guinea, West Africa [1] in December 2013, and over two years later sporadic cases were still occurring. The outbreak spread to multiple countries with Guinea, Sierra Leone and Liberia being particularly affected. As of January, over 28,600 cases and more than 11,300 deaths have been reported by the World Health Organization (WHO) [2]. In addition to cases in West Africa, infected personnel have been evacuated out of Africa, and facilities in Europe and the United States have treated Ebola virus (EBOV)-infected patients. An unprecedented international response led to volunteers from all over the world travelling to West Africa to work in laboratories, hospitals, treatment centres, holding centres or in the local communities to provide diagnostics, treatment and care and advice to try and halt the spread of the disease. Often working in temporary or adapted buildings with re-purposed equipment, personnel were reliant on personal protective equipment and disinfection for effective infection control.

EBOV, along with the other filoviruses, is a lipid enveloped virus, often considered fragile, that can be inactivated by the use of both physical and chemical methods [3]. Until recently there was limited data on efficacy of disinfectants against EBOV. The Center for Disease Control (CDC), advises “Suitable disinfectant solutions include 0.5% sodium hypochlorite (10% aqueous solution of household bleach), as well as fresh, correctly prepared solutions of glutaraldehyde (2% or as recommended by the manufacturer) and phenolic disinfectants (0.5%–3%)” [4]. The use of sodium hypochlorite solutions
is widely advised in further CDC guidance on managing suspected viral hemorrhagic fever [5] and
by the WHO in their guidelines on laboratory diagnosis of EVD [6]. Many chlorine-based products,
however, are corrosive and may damage fragile equipment or apparatus that needs to be re-used [7].
Chlorine-based products may compromise performance on vehicles used for transport of infected
patients, including military aircraft, which were used for evacuation of infected personnel to Europe
and the United States. Therefore, alternative disinfectants that are effective against EBOV need to be
identified and tested.

Two commonly used military detergents (Calla 1452 and Ardrox 6092) were assessed in a contact
kill assay for efficacy against a single EBOV strain in comparison with sodium hypochlorite,
and the commercially available disinfectant Desintex. Efficacy was tested on two representative
military-relevant surfaces: painted aluminium as a hard, non-porous material, and pilot seat-belt
strapping material as a soft porous material. British standard requirements for virucidal activity state
a product should “demonstrate at least a decimal log reduction of 4 in virus titre” when tested in
accordance with various parameters [8].

2. Materials and Methods

2.1. Test Materials

Tissue culture media (TCM) used throughout was Dulbecco’s minimum essential media
supplemented with 2% foetal calf serum, 1% L-glutamine and 1% penicillin/streptomycin (all Sigma,
Gillingham, Dorset, UK). The fifty percent tissue culture infectious dose (TCID\text{50}) amount was
calculated using the method of Reed and Muench [9].

EBOV H.sapiens-tc/COD/1976/Yambuku-Ecran, hereafter referred to as EBOV-Ecran was used in
all studies. This virus, previously known as EBOV “E718” was supplied by Public Health England [10].
Passage 5 material was used to infect Vero C1008 cells (ECACC Cat. No. 85020206) maintained in
TCM. Virus was harvested on day 5 post-inoculation and titrated to produce a working stock at
1 \times 10^7 \text{TCID}_\text{50}/\text{mL}. EBOV-Ecran was titrated in 96-well plates using the endpoint \text{TCID}_\text{50} assay as
previously described [11], although modified with removal of the staining and fixing steps so that all
plates were scored for cytopathic effects upon microscopic examination on day 7.

Four disinfectants were assessed for efficacy against EBOV-Ecran (Table 1). Calla 1452 (Zip-chem
products, Morgan Hill, CA, USA) was prepared by dilution in tap water (1:32) to give an approximate
starting concentration of 3%, as recommended by Air Movements Navy Command Headquarters.
Desintex (Laboratoires Rochex, Ville La Grand, France) was prepared by dilution in tap water to
a starting concentration of 5\% v/v. Ardrox 6092 (Chemetall Plc, Bletchley, Milton Keynes, UK Defence
Standard 79-17/2) required no preparation as this was used as a neat solution, and sodium hypochlorite
(SyChem, UK) was prepared by dilution in tap water to a starting concentration of 1.5\% v/v.

Aircraft relevant materials were supplied directly by 1710 Naval Air Squadron as pre-cut coupons
and were not pre-conditioned in any way. The coupons assessed were painted aluminium and pilot
seat-belt strapping material in 1 cm\textsuperscript{2} discs.

All growth and manipulation of EBOV-Ecran was performed within a Class III microbiological
safety cabinet within a dedicated containment level 4 laboratory at Defence Science and Technology
Laboratory (Dstl), Porton Down [12].
| Name          | Supplier                  | Active Components (%)                                                                 | Preparation          | Final % Tested (v/v) | Volume Tested (µL) | Contact Time (Mins at RT) |
|---------------|---------------------------|----------------------------------------------------------------------------------------|-----------------------|----------------------|---------------------|--------------------------|
| Ardrox 6092   | Chemetall, UK             | 3-Butoxypropan-2-ol, propylene glycol monobutyl ether (3–5); Disodium metasilicate (1–2.5); Alcohol ethoxylate (1–2.5) | Used neat            | 50.0                 | 50                  | 10                       |
| Calla 1452    | Zip-chem products, CA, US | Alkyl Dimethyl benzylammonium Chloride (1.1); Di(octyl-decyl) dimethyl ammonium Chloride (1.6); Ethanol (0.3) | 1:32 dilution in tap water | 1.5                  | 50                  | 10                       |
| Desintex      | Laboratoires Rochex, France | Alkylamine (8.7); PolyHexaMethyleneBiguanide (2.1); Quaternary ammonium (3.1)         | 1:20 dilution in tap water | 2.5                  | 50                  | 10                       |
| Sodium Hypochlorite | SyChem, UK               | Sodium hypochlorite (14–15); Potassium permanganate (<1)                                | 1:10 dilution in tap water | 0.75                 | 50                  | 10                       |

2.2. Recovery Testing

To optimise recovery of EBOV-Ecran from different materials, virus was deposited onto coupons of aluminium or strapping in 50 µL droplets at room temperature. Samples were then placed into TCM sufficient to be submerged (2 mL), and vortexed for 10 s before being enumerated by standard TCID₅₀ assay. Recovery of EBOV-Ecran from sterile 6-well cell culture plates served as a positive control and recovery of TCM only from aluminium and strapping was used as a negative control. All recovery testing was performed in triplicate. In an attempt to optimise recovery, vortex times were increased or samples were vortexed multiple times.

2.3. Toxicity Testing

To test the toxicity of disinfectants against the Vero C1008 cell line, each disinfectant was added to a contact plate and incubated at room temperature for 10 min. Samples were then washed by centrifugation at 10,000 rpm for 5 min in a Microcentaur MSE centrifuge allowing the formation of a pellet which was re-suspended in TCM and enumerated using the TCID₅₀ assay. Cells were observed for signs of toxicity. EBOV-Ecran only (no disinfectant) was used as a positive control, and TCM without virus was used as a negative control. All testing was performed in triplicate. To reduce toxicity, the number of washes was increased as necessary.

2.4. Efficacy of Different Disinfectants by Virus Enumeration

Coupon assessments were made with a single contact time of 10 min at room temperature (Table 1). In triplicate, 50 µL of disinfectant (or TCM as a positive control) was added to 50 µL EBOV-Ecran on aluminium or strapping coupons, or directly to a sterile 6-well cell culture plate. Additionally, disinfectant was added to 50 µL TCM only as a negative control. After 10 min contact time, virus was recovered by vortexing for 10 s followed by washing by centrifugation with TCM up to three times, sufficient to reduce toxicity to Vero C1008 cells. TCID₅₀ enumeration was performed on all samples. Test samples were evaluated against the control samples with a 4-log₁₀ reduction in viral titre being the standard minimum for a disinfectant to be considered effective.

2.5. Efficacy of Different Disinfectants by Cell Passage

Samples were also qualitatively assessed for inactivation by serial passage [13]. Washed samples generated as above were used to infect flasks of Vero C1008 cells (5 mL total volume) and incubated at 37 °C/5% CO₂ for 7 days. After this time, flasks were observed for signs of infection (cytopathic effects, CPE) and the flasks were passaged in to a larger volume (10 mL) of fresh TCM and incubated for a further 7 days. After observation for CPE, flasks were passaged a third and final time into 20 mL TCM and incubated for another 7 days. If no signs of infection (no CPE) were observed after 3 passages, samples were considered to be free of viable virus.
2.6. Statistical Analysis

GraphPad Prism (version 6.02, La Jolla, CA, USA) was used to analyse the efficacy data. Two-way ANOVA with Bonferroni’s multiple comparison test was used for each coupon type to compare the mean log reduction in titre from control of each disinfectant to the log reduction when no disinfectant was used (0) at the 95% confidence level. Additionally, identical 2-way ANOVA analysis was used for each disinfectant to compare the mean log reduction in titre of the different coupon types.

3. Results

3.1. Recovery of EBOV-Ecran from Test Surfaces and Toxicity of Disinfectants to Cells

In triplicate, samples of EBOV-Ecran were deposited on two test surfaces, aluminium and pilot strapping material. Results are shown in Figure 1. The mean titre of EBOV-Ecran recovered from a sterile cell culture plate (no coupon) was $1.3 \times 10^6$ TCID$_{50}$/mL. The mean titre recovered from aluminium coupons was $5.9 \times 10^5$ TCID$_{50}$/mL, representing a 45% recovery efficiency and a high enough titre to determine a 4-log$_{10}$ reduction. No further attempts were made to optimise recovery from aluminium coupons. The mean titre recovered from strapping was $4.1 \times 10^5$ TCID$_{50}$/mL. This represents a recovery efficiency of only 0.32%. The limit of quantification of the TCID$_{50}$ assay is 10 TCID$_{50}$/mL, therefore a starting titre of at least $1 \times 10^5$ TCID$_{50}$/mL is required to demonstrate a 4-log$_{10}$ reduction. Attempts to improve recovery from strapping were not successful (results not shown). Therefore, in addition to reduction in viral titre, strapping samples were subjected to serial passage in Vero C1008 cells to confirm inactivation.

![Figure 1. Recovery of Ebola virus (EBOV)-Ecran from different test surfaces. EBOV-Ecran or tissue culture media (TCM) was deposited onto 2 different coupon types, or directly onto a sterile 6-well cell culture plate (no coupon), before being recovered into 2 mL TCM by vortexing for 10 s. Samples were washed up to three times prior to enumeration. The titre of EBOV-Ecran (in TCID$_{50}$/mL) recovered from 3 replicate tests is shown (±SEM). All TCM samples were negative for viable EBOV-Ecran. TCID$_{50}$: 50% tissue culture infectious dose.](image)

Four disinfectants (see Table 1) were tested for toxicity in cells in triplicate. Calla 1452, Ardrox 6092 and sodium hypochlorite were rendered non-toxic to cells after one wash in TCM (results not shown). Desintex required three washes to remove toxic components (results not shown).
3.2. **Coupon Disinfectant Assays**

Four disinfectants (see Table 1) were tested for the ability to reduce the viral titre of EBOV-Ecran on different coupon types. All samples were tested by enumeration in TCID\(_{50}\)/mL after a 10 min contact time at room temperature (Table 2). The limit of quantification (LoQ) of the TCID\(_{50}\) assay is 10 TCID\(_{50}\)/mL. This equates to 50% (4/8 well in a 96-well plate format) of neat sample being observed to have CPE in the TCID\(_{50}\) assay. However, <50% of the inoculated wells may be positive for CPE (e.g., 1, 2 or 3 of 8 positive wells in the neat dilution), indicating viable virus is present but is <10 TCID\(_{50}\)/mL. This number cannot truly be enumerated by the method of Reed and Muench [9].

### Table 2. Recovery of EBOV-Ecran recovered from different test surfaces after treatment with four disinfectants.

| Sample Type                        | Mean Titre of EBOV-Ecran Recovered (TCID\(_{50}\)/mL) |
|------------------------------------|------------------------------------------------------|
|                                    | Aluminium \((n = 3)\)  | Strapping \((n = 3)\)  | No Coupon \((n = 6)\) |
| EBOV-Ecran + TCM                   | \(1.5 \times 10^3\)  | \(2.2 \times 10^2\)  | \(1.1 \times 10^3\) |
| EBOV-Ecran + Ardrox 6092           | 0*                     | 0*                     | 0*                     |
| EBOV-Ecran + Calla 1452            | \(7.0 \times 10^2\)  | \(2.3 \times 10^2\)  | 7.0**                  |
| EBOV-Ecran + sodium hypochlorite    | 0*                     | 0*                     | 0*                     |
| EBOV-Ecran + Desintex              | \(1.6 \times 10^1\)  | 0*                     | 2.4**                  |

TCID\(_{50}\)/mL: 50% tissue culture infectious dose per mL; * No viable virus observed in TCID\(_{50}\) assay; ** Extrapolated, virus observed but below limit of quantification (LoQ) (10 TCID\(_{50}\)).

Efficacy of the four disinfectants was determined by comparing the mean log reduction in EBOV-Ecran with the positive control (EBOV-Ecran + TCM; Table 2, Figure 2). The British standards require a 4-log\(_{10}\) reduction of virus for a disinfectant to be classed as effective [8].

![Figure 2](image-url)  
*Figure 2.* Reduction in EBOV-Ecran titre on different test surfaces after contact with four different disinfectants. EBOV-Ecran was deposited onto three surfaces and an equal volume of disinfectant or TCM was added for a contact time of 10 min \((n = 3)\). When TCM was added to virus in place of disinfectant, high titres of EBOV-Ecran were recovered (Table 2; \(1.5 \times 10^3\) TCID\(_{50}\)/mL from aluminium; \(2.2 \times 10^3\) TCID\(_{50}\)/mL from strapping and \(1.1 \times 10^5\) TCID\(_{50}\)/mL from no coupon) and these values served as the positive control against which disinfectants were assessed. The reduction in log\(_{10}\) titre was determined as the difference between the mean log\(_{10}\)-titre with no disinfection (TCM only) and the mean log\(_{10}\)-titre for each disinfectant on aluminium coupons (grey bars), strapping coupons (black bars) or directly onto a sterile 6-well cell culture plate (no coupon, white bars). The British standard minimum requirement is a 4-log\(_{10}\) reduction (dashed line).
3.2.1. Efficacy of Four Disinfectants against EBOV-Ecran on Aluminium Coupons

For the aluminium coupons, under the parameters tested, the mean titre of EBOV-Ecran recovered from the coupon was \(1.5 \times 10^5\) TCID\(_{50}\)/mL. Addition of sodium hypochlorite or Ardrox 6092 resulted in no detectable virus in the TCID\(_{50}\) assay (Table 2). There was also no detectable EBOV-Ecran when Ardrox 6092 and sodium hypochlorite were added to virus on a sterile cell culture plate (no coupon; Table 2). When Calla 1452 was added to EBOV-Ecran on aluminium coupons \(7 \times 10^2\) TCID\(_{50}\)/mL of EBOV-Ecran was recovered from the sample and when Calla 1452 and EBOV-Ecran were mixed on a sterile cell culture plate, viable virus was also seen (Table 2). Low levels of EBOV-Ecran were detected when Desintex was added to aluminium coupons and viable virus was also observed, albeit below the LoQ of the assay, when Desintex and EBOV-Ecran were tested on a sterile cell culture plate only (Table 2).

For the aluminium coupons, a greater than 4-log\(_{10}\) drop in EBOV-Ecran titre was observed with Ardrox 6092 and sodium hypochlorite (Figure 2). Based on mean counts, a 4-log\(_{10}\) drop in EBOV-Ecran titre was also observed with Desintex on aluminium, although viable virus was recovered (Table 2). When no coupon was present, a 4-log\(_{10}\) drop in EBOV-Ecran titre was observed for all four disinfectants (Figure 2).

3.2.2. Efficacy of Four Disinfectants against EBOV-Ecran on Strapping Coupons

Poor recovery was observed for pilot strapping (Figure 1), with a mean titre of \(2 \times 10^3\) TCID\(_{50}\)/mL. A 4-log\(_{10}\) reduction in titre therefore fell below the LoQ of the assay. Samples were still processed in the TCID\(_{50}\) assay despite the inability to observe the necessary reduction in order to identify a degree of activity prior to serial cell passage. As with the aluminium coupons, no viable virus could be detected on strapping coupons treated with Ardrox 6092 or sodium hypochlorite in the TCID\(_{50}\) assay (Table 2). When Calla 1452 was added to EBOV-Ecran on strapping, \(2.3 \times 10^2\) TCID\(_{50}\)/mL of EBOV-Ecran was recovered and when Desintex and EBOV-Ecran were tested on strapping coupons no viable virus was detected (Table 2). The use of TCM in place of EBOV-Ecran as a negative control was conclusive for all samples types as no viable virus was detected (results not shown). Due to poor recovery, a 4-log\(_{10}\) reduction in titre was not achieved with any disinfectant on the strapping coupons (Figure 2).

Additional qualitative testing of disinfectants on strapping coupons was performed by serial passage. All samples from strapping coupons treated with disinfectant or media only were subject to three passages in cells to allow for amplification of low numbers of viable virus to detectable limits. After three passages, no viable virus was observed in replicate Ardrox 6092- or sodium hypochlorite-treated samples suggesting there had been complete inactivation of EBOV-Ecran on the strapping coupons when treated with these disinfectants. Desintex-treated strapping coupons were also all negative for viable virus after three passages, with the exception of one of the Desintex-treated EBOV-Ecran samples from a sterile cell culture plate (no coupon) after the second passage. This suggests that this concentration of Desintex does not consistently provide complete inactivation. Calla 1452-treated samples were positive for viable virus after the first passage (when recovered from strapping coupons and after the second passage when recovered from a sterile cell culture plate (no coupon).

Two-way ANOVA analysis with Bonferroni’s multiple comparison test indicated that the difference in log reduction for each disinfectant compared to the log reduction with no disinfectant (TCM only, 0, no reduction) was significant for Ardrox 6092, sodium hypochlorite and Desintex on all three surface types (\(p \leq 0.01\) for aluminium and for when no coupon was used and \(p \leq 0.05\) for strapping). Calla 1452 produced a significant log reduction from control only when no coupon was present (\(p \leq 0.01\)). For each disinfectant, there was no significant difference in log reduction between the three different surface types.
4. Discussion and Conclusions

The worst outbreak of EVD in history has highlighted the need for effective disinfectants [1,2,4–6]. Although efficacy is the primary objective, the disinfectants also need to be appropriate for their intended use. Disinfectants may be required for use on the skin and body parts, complex surfaces, delicate instruments and textiles and on small or large areas. Whilst chlorine-based products are widely recommended for use against EBOV, and used internationally in hospitals, laboratories and in the community, they can also be damaging. Equipment or apparatus that is damaged may lose integrity, utility or reduce safe operation, therefore alternative effective disinfectants are needed. Three disinfectants for which there is no published data available were tested for activity against EBOV-Ecran alongside sodium hypochlorite as a control disinfectant on two representative aircraft relevant surfaces. The surfaces, painted aluminium and pilot seat-belt strapping, were also representative of other porous or non-porous surfaces that may be encountered on transport vehicles, buildings or other settings.

As perhaps expected, sodium hypochlorite performed well in our assessments in that no viable virus was recovered from either aluminium or strapping coupons, or when no coupon was present. Different chlorine-based products will have different properties and will react differently with different material. A single concentration of sodium hypochlorite (SyChem) was tested here, but 0.75% sodium hypochlorite met the minimum British standards under the conditions tested. The efficacy of sodium hypochlorite against a different EBOV has already been investigated by others using similar methods to our work [14]. In this work, 0.5% and 1.0% sodium hypochlorite gave complete reduction in viral titre after 5 min contact time whereas concentrations of 0.1% and 0.01% were not effective in reducing viral titre over 10 min contact time [14]. Our data is consistent with these results and provides evidence of a further concentration of sodium hypochlorite (0.75%) demonstrating efficacy against EBOV-Ecran. For General Healthcare Settings in West Africa the CDC recommends 0.5% as strong for disinfecting ‘surfaces, objects, medical equipment, and gloved hands’ and a milder 0.05% for hand washing etc. [15]. The lower concentration may not be effective but higher concentrations have been reported to affect equipment [7] and personnel [16] and were shown to be damaging to aircraft surfaces [17]. An alternative to liquid chlorine based products for general disinfection is the use of gaseous chlorine dioxide which has been used to decontaminate hospitals and large areas previously [18,19]. However, surrogate testing suggests this method should not be relied on solely to be effective against EBOV [20] and further efficacy testing and assessment for corrosiveness and potential damage from this form of disinfection is required. Likewise, other forms of gaseous disinfection (e.g., hydrogen peroxide) or other methods of inactivation (such as UV light) could be considered. However, in an outbreak environment where there may be limited infrastructure, training or there may not be the appropriate personal protective equipment to work with more novel disinfection methods, simpler methods are likely to be more advantageous.

The active components of Desintex are alkylamine and quarternary ammonium compounds. Our results demonstrated good efficacy of 2.5% Desintex although not complete inactivation. A reduction in titre of EBOV-Ecran by over 3-log₁₀ was observed on both coupon types as well as on sterile cell culture plates. However, viable virus was detected when Desintex was added to EBOV-Ecran on aluminium coupons and when no coupon was present. EBOV-Ecran was not detected on strapping coupons although recovery levels were low from this coupon type, and when samples were serially passaged there was no detectable virus-induced CPE. Although under the conditions tested Desintex meets the British Standard requirements, this is a minimum requirement and the fact that viable virus remained should be noted. EBOV strains have been shown to be infectious at low levels, and to remain infectious in different matrices [21–24]. Desintex has also been shown to be damaging to aircraft surfaces [17] so may not be a suitable alternative to sodium hypochlorite for use on aircraft. It is also important to note that Dstl routinely uses Desintex at a final concentration of 5% v/v, twice the concentration tested in these studies.
Ardrox 6092 is a water-based alkaline (pH > 11) cleaner designated as suitable for cleaning civil and military fixed wing aircraft and helicopters. Calla products contain ethanol, alkyl dimethyl benzyl- and di(octyl-decyl)-ammonium chloride and are also sold as aircraft cleaners. Calla 1452 has been included in the cleaning supplies on aircraft used to move EBOV-infected patients [25] as it is specifically a hard surface, interior aircraft disinfectant. Both Ardrox 6092 and Calla 1452 are commonly used by the UK military for routine cleaning of aircraft and do not damage the airframes [17] but their efficacy against EBOV had not been determined. Our results show that Calla 1452 is not effective against EBOV-Ecran on the two coupons tested; under the conditions described Calla 1452 showed minimal reduction in viral load falling below the level required to meet British disinfection standards [8]. However, the efficacy of Calla 1452 may be improved with increased contact times or different concentrations. The results generated in this study would suggest that it is not currently advisable to use Calla 1452 to disinfect aircraft surfaces that may be contaminated with EBOV.

Ardrox 6092 was demonstrated to show efficacy against EBOV-Ecran under the conditions tested and therefore further efficacy characterisation should be pursued for example different contact times, disinfectant concentrations or volumes or different matrices. Ardrox 6092 is known to be non-corrosive to aircraft [17] but the effect of high pH on virus activity had not been determined. Viable virus was not recovered from aluminium or strapping coupons when Ardrox 6092 was applied although it was not possible to determine a 4-log reduction for the strapping coupons (a 3-log reduction was achieved). Recovery from strapping was poor; despite numerous attempts to increase recovery, and success with other porous material, it was not possible to improve efficiency from strapping. Poor recovery from airline carpet material had been previously observed [26] whilst a recent study using providone-iodine hand wash solutions against EBOV described an alternative method of large volume plating to verify efficacy and increased sensitivity when low titres of stock EBOV were harvested from culture [27]. Utilising this method may have improved sensitivity when assaying strapping samples but we did not have the capability to do this or to produce a higher titre starting stock. The promising efficacy of Ardrox 6092 against EBOV-Ecran necessitates inclusion in future studies and for further assessment against other viruses, surfaces and matrices.

Further assessments of disinfectant action on strapping material were made by serial cell passage in order to amplify any low levels of virus that may have been present in the sample and therefore increased the sensitivity of the testing. Previously it has been shown that 0.1 TCID<sub>50</sub> can be amplified by passage in tissue culture flasks [13], therefore the results using Ardrox 6092 and the detection limit of the flask passage under the conditions tested in these experiments, indicated that Ardrox 6092 is likely to meet the minimum standards required. However, improved recovery efficiency or an alternative method of analysis would be required to determine this.

Our studies were limited to testing EBOV-Ecran only in TCM which is not representative of the type of matrix that may be encountered in the field, where EBOV will be present in bodily fluids including but not limited to, blood, vomit and faeces [26]. The survival of different isolates of EBOV in blood [14,21,22,24,26] and in other bodily fluids or specimens [24,26] has been reported. The organic matter in more complex matrices such as bodily fluids may initiate a reaction with the disinfectant or may provide protection to the virus, affecting the mode of action of disinfectant. Complications may also arise from potential sample toxicity and recovery of virus thereby rendering results more difficult to interpret. TCM is a representative matrix encountered in the laboratory and provides an initial starting point in the assessment of disinfectant efficacy against pathogenic viruses. Disinfectants that show efficacy in TCM can be taken forward for further study in more complex matrices, whilst disinfectants that do not work in TCM are un-likely to be efficacious in more complex media although further testing in alternative matrices is recommended.

Additionally, we did not use the EBOV strain responsible for the West Africa outbreak (EBOV-Makona) due to availability at the time, lack of characterisation and low titre during initial experiments. However, initial phylogenetic analysis of isolates from West Africa shows they are a distinct clade but still have high homology with early EBOV strains [1,28]. The protocols performed
in these studies may be considered a ‘worse-case scenario’ as there was no mixing of virus and disinfectant, and equal volumes were used. There are multiple factors and conditions that contribute to a disinfectant’s effectiveness and it was not in the scope of this study to investigate all parameters. Further work is required to fully characterize the efficacy of these and other disinfectants for use against highly pathogenic viruses.

The need to find further disinfectants that are effective against EBOV that can be used in a range of scenarios remains. Under the conditions tested in this study, Calla 1452 has limited efficacy against EBOV-Ecran, but aircraft cleaner Ardrox 6092 may be an effective alternative to corrosive sodium hypochlorite for aircraft relevant materials and other vulnerable surfaces.

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References
1. Baize, S.; Pannetier, D.; Oestereich, L.; Rieger, T.; Koivogui, L.; Magassouba, N.; Soropogui, B.; Sow, M.S.; Keita, S.; De Clerck, H.; et al. Emergence of Zaire Ebola virus disease in Guinea. N. Engl. J. Med. 2014, 371, 1418–1425. [CrossRef] [PubMed]
2. World Health Organization Situation Reports. Available online: http://www.who.int/csr/disease/ebola/situation-reports/archive/en/ (accessed on 31 May 2016).
3. Mitchell, S.W.; McCormick, J.B. Physicochemical inactivation of Lassa, Ebola, and Marburg viruses and effect on clinical laboratory analyses. J. Clin. Microbiol. 1984, 20, 486–489. [PubMed]
4. CDC. Management of Patients with Suspected Viral Hemorrhagic Fever. Available online: http://www.cdc.gov/mmwr/preview/mmwrhtml/00037085.htm (accessed on 21 April 2016).
5. CDC. Interim Guidance for Managing Patients with Suspected Viral Hemorrhagic Fever in U.S. Hospitals. Available online: http://www.cdc.gov/HAI/pdfs/bbp/VHFinterimGuidance05_19_05.pdf (accessed on 21 April 2016).
6. WHO. Laboratory Diagnosis of Ebola Virus Disease. Available online: http://apps.who.int/iris/bitstream/10665/134009/1/WHO_EVD_GUIDANCELAB_14.1_eng.pdf (accessed on 21 April 2016).
7. Johnston, A.M.; Lewis, S.E. Decontaminating Ebola-infected ultrasound probes. Anaesthesia 2015, 70, 628–629. [CrossRef] [PubMed]
8. British Standards Institution. BSEN 14476:2013+A1:2015. Chemical Disinfectants and Antiseptics. Quantitative Suspension Test for the Evaluation of Virucidal Activity in the Medical Area. Test Method and Requirements (Phase 2/Step 1). 2013. Available online: http://shop.bsigroup.com/ProductDetail/?pid=000000000030318020 (accessed on 9 July 2015).
9. Reed, L.J.; Muench, H. A simple method of estimating fifty per cent endpoints. Am. J. Hyg. 1938, 27, 493–497.
10. Kuhn, J.H.; Lofts, L.L.; Kugelman, J.R.; Smither, S.J.; Lever, M.S.; van der Groen, G.; Johnson, K.M.; Radoszitzky, S.R.; Bavari, S.; Jahrling, P.B.; et al. Reidentification of Ebola virus E718 and ME as Ebola Virus/H.sapiens-tc/COD/1976/Yambuku-Ecran. Genome Announc. 2014, 2, e1178–e1114. [CrossRef] [PubMed]
11. Smither, S.J.; Lear-Rooney, C.; Biggins, J.; Pettitt, J.; Lever, M.S.; Olinger, G.G., Jr. Comparison of the plaque assay and 50% tissue culture infectious dose assay as methods for measuring filovirus infectivity. J. Virol. Methods 2013, 193, 565–571. [CrossRef] [PubMed]
12. Smither, S.J.; Lever, M.S. A review of filovirus work and facilities at the Defence Science and Technology Laboratory Porton Down. Viruses 2012, 4, 1305–1317. [CrossRef] [PubMed]
13. Smither, S.J.; Weller, S.A.; Phelps, A.; Eastaugh, L.; Ngugi, S.; O’Brien, L.M.; Steward, J.; Lonsdale, S.G.; Lever, M.S. Buffer AVL alone does not inactivate Ebola virus in a representative clinical sample type. *J. Clin. Microbiol.* 2015, 53, 3148–3154. [CrossRef] [PubMed]

14. Cook, B.W.; Cutts, T.A.; Nikiforuk, A.M.; Poliquin, P.G.; Court, D.A.; Strong, J.E.; Theriault, S.S. Evaluating environmental persistence and disinfection of the Ebola virus Makona variant. *Viruses* 2015, 7, 1975–1986. [CrossRef] [PubMed]

15. CDC. For General Healthcare Settings in West Africa: How to Prepare and Use Chlorine Solutions. Available online: http://www.cdc.gov/vhf/ebola/hcp/mixing-chlorine-solutions.html (accessed on 31 May 2016).

16. Carpenter, A.; Cox, A.T.; Marion, D.; Phillips, A.; Ewington, I. A case of a chlorine inhalation injury in an Ebola treatment unit. *J. R. Army Med. Corps* 2015, 162, 229–231. [CrossRef] [PubMed]

17. Andrew Dutch, NAVY 1710NAS-MIG CCHS 2. Personal communication.

18. Lowe, J.J.; Gibbs, S.G.; Iwen, P.C.; Smith, P.W.; Hewlett, A.L. Decontamination of a hospital room using gaseous chlorine dioxide: *Bacillus anthracis*, *Francisella tularensis*, and *Yersinia pestis*. *J. Occup. Environ. Hyg.* 2013, 10, 533–539. [CrossRef] [PubMed]

19. Doona, C.J.; Feeherry, F.E.; Kustin, K.; Olinger, G.G.; Setlow, P.; Malkin, A.J.; Leighton, T. Fighting Ebola with novel spore decontamination technologies for the military. *Front. Microbiol.* 2015, 6, 663. [CrossRef] [PubMed]

20. Lowe, J.J.; Hewlett, A.L.; Iwen, P.C.; Smith, P.W.; Gibbs, S.G. Surrogate testing suggests that chlorine dioxide gas exposure would not inactivate Ebola virus contained in environmental blood contamination. *J. Occup. Environ. Hyg.* 2015, 8. [Epub ahead of print]. [CrossRef] [PubMed]

21. Piercy, T.J.; Smither, S.J.; Steward, J.A.; Eastaugh, L.; Lever, M.S. The survival of filoviruses in liquids, on solid substrates and in a dynamic aerosol. *J. Appl. Microbiol.* 2010, 109, 1531–1539. [CrossRef] [PubMed]

22. Fischer, R.; Judson, S.; Miazgowicz, K.; Bushmaker, T.; Prescott, J.; Munster, V.J. Ebola virus stability on surfaces and in fluids in simulated outbreak environments. *Emerg. Infect. Dis.* 2015, 2, 1243–1246. [CrossRef] [PubMed]

23. Judson, S.; Prescott, J.; Munster, V. Understanding ebola virus transmission. *Viruses* 2015, 7, 511–521. [CrossRef] [PubMed]

24. Prescott, J.; Bushmaker, T.; Fischer, R.; Miazgowicz, K.; Judson, S.; Munster, V.J. Postmortem stability of Ebola virus. *Emerg. Infect. Dis.* 2015, 21, 856–859. [CrossRef] [PubMed]

25. Thoms, W.E.; Wilson, W.T.; Grimm, K.; Conger, N.G.; Gonzales, C.G.; DeDecker, L.; Hatzfeld, J.J. Long-range transportation of Ebola-exposed patients: An evidence-based protocol. *Am. J. Infect. Dis. Microb.* 2015, 2, 19–24.

26. Schuit, M.; Miller, D.M.; Reddick-Elick, M.S.; Wlazlowski, C.B.; Filone, C.M.; Herzog, A.; Colf, L.A.; Wahl-Jensen, V.; Hevey, M.; Noah, J.W. Differences in the comparative stability of Ebola virus Makona-C05 and Yambuku-Mayinga in Blood. *PLoS ONE* 2016, 11, e0148476. [CrossRef] [PubMed]

27. Eggers, M.; Eickmann, M.; Kowalski, K.; Zorn, J.; Reimer, K. Povidone-iodine hand wash and hand rub products demonstrated excellent in vitro virucidal efficacy against Ebola virus and modified vaccinia virus Ankara, the new European test virus for enveloped viruses. *BMC Infect. Dis.* 2015, 15, 375. [CrossRef] [PubMed]

28. Gire, S.K.; Goba, A.; Andersen, K.G.; Sealfon, R.S.; Park, D.J.; Kanneh, L.; Jalloh, S.; Momoh, M.; Fullah, M.; Dudas, G.; et al. Genomic surveillance elucidates Ebola virus origin and transmission during the 2014 outbreak. *Science* 2014, 345, 1369–1372. [CrossRef] [PubMed]