# Marvel analysis of the measured high-resolution rovibrational spectra of $\mathrm{H}_{2}{ }^{32}$ S 

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#### Abstract

44325 measured and assigned transitions of $\mathrm{H}_{2}{ }^{32}$ S, the parent isotopologue of the hydrogen sulfide molecule, are collated from 33 publications into a single database and reviewed critically. Based on this information, rotation-vibration energy levels are determined for the ground electronic state using the Measured Active Rotational-Vibrational Energy Levels (MARVEL) technique. The ortho and para principal components of the measured spectroscopic network of $\mathrm{H}_{2}{ }^{32} \mathrm{~S}$ are considered separately. The verified set of 25293 ortho- and 18778 para- $\mathrm{H}_{2}{ }^{32} \mathrm{~S}$ transitions determine 3969 ortho and 3467 para energy levels. The Marvel results are compared with alternative data compilations, including a theoretical variational linelist.


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## 1. Introduction

Hydrogen sulfide, $\mathrm{H}_{2} \mathrm{~S}$, is a volcanic gas present on earth and other geologically active planets and moons, such as Io [1], Venus [2], and, theoretically, hot super-earth "lava" exoplanets [3]. It has recently been detected above the clouds in the atmosphere of Uranus [4]. On earth, microbial respiration of seawater sulfate creates a hydrogen sulfide-rich ecosystem [5], suggesting that $\mathrm{H}_{2} \mathrm{~S}$ is a component for a potential microbial life sustaining atmosphere on Venus [2], and a potential biomarker for life on exoplanets [6]. $\mathrm{H}_{2} \mathrm{~S}$ influences many physiological processes [7], it is of importance in the treatment of respiratory diseases [8], and it is used as a measure of the quality of air near oil refineries [9]. The parent isotopologue, $\mathrm{H}_{2}^{32} \mathrm{~S}$, was first detected in interstellar space in 1972 [10]. Many of the scientific and engineering applications mentioned re-

[^0]quire the detailed knowledge of the rovibrational energy levels of $\mathrm{H}_{2}{ }^{32} \mathrm{~S}$.

As part of this work, we present the largest compilation of published experimental rovibrational transition data for $\mathrm{H}_{2}^{32} \mathrm{~S}$. The experimental database of $\mathrm{H}_{2}^{32} \mathrm{~S}$ transitions has been formatted and analysed using the Marvel (Measured Active Rotational-Vibrational Energy Levels) spectroscopic network (SN) software [11-13]. This study builds on the data of the 2012W@DIS information system for the hydrogen sulfide molecule [14].

As to the structure of this paper, the next section provides the theory underlying the present study. Section 3 presents and discusses the experimental sources used, with results given in Section 4. Section 5 discusses these results and compares the empirical rovibrational energies presented in this work with corresponding levels previously determined both by experiment and theory. Finally, Section 6 provides our conclusions. All transition data and energy levels resulting from this work are included as supplementary data files.

## 2. Theory

### 2.1. MARVEL

The Marvel procedure [11-13] is based on the theory of spectroscopic networks [15,16], with the energy levels represented as nodes and the transitions between them as edges. The related Marvel code can be used to critically evaluate and validate experimentally-determined transition wavenumbers and uncertainties collected from the literature, inverting the wavenumber information to obtain accurate empirical energy levels with an associated uncertainty. Marvel has been successfully used to evaluate the energy levels for molecules such as ${ }^{12} \mathrm{C}_{2}$ [17], ${ }^{48} \mathrm{Ti}{ }^{16} \mathrm{O}$ [18], water vapour [19-23], $\mathrm{H}_{3}^{+}$[24], $\mathrm{H}_{2} \mathrm{D}^{+}$and $\mathrm{D}_{2} \mathrm{H}^{+}$[25], ${ }^{12} \mathrm{C}_{2} \mathrm{H}_{2}$ [26], ${ }^{14} \mathrm{NH}_{3}$ [27,28], and ${ }^{12} \mathrm{C}_{2} \mathrm{H}_{2}^{16} \mathrm{O}$ [29]. Marvel requires each measured transition to have an associated uncertainty and for each energy level considered to possess a unique set of quantum numbers.

### 2.2. Quantum number labelling

The seven quantum numbers that were used for labelling the upper and lower rovibrational states of $\mathrm{H}_{2}^{32} \mathrm{~S}$ are the same as those used in a previous Marvel investigation to classify water [19]. These quantum numbers are summarised in Table 1. Normal-mode labelling is used for the vibrations, where $\nu_{1}, \nu_{2}$ and $\nu_{3}$ stand for the symmetric stretch, bend, and antisymmetric stretch vibrations, respectively. Standard asymmetric-top quantum numbers are used for the rotations, where $J, K_{a}$, and $K_{c}$ are the three quantum numbers associated with rotational angular momentum, $\mathbf{J}$, and the two projections along the $A$ and $C$ axes. We also provide, as part of the label, the nuclear spin state (ortho or para), which is deduced by whether ( $v_{3}+K_{a}+K_{c}$ ) is odd (ortho) or even (para) [26,30]. Hyperfine coupling associated with nuclear spins has been neglected.

When local-mode notation was used by an experimental source considered in this work, the quantum numbers were transformed to the normal-mode notation ( $v_{1} v_{2} v_{3}$ ) to describe a vibrational state. Note that some of the data sources considered in this study use $J$ and $\tau=K_{a}-K_{c}$ instead of $J, K_{a}$, and $K_{c}$ [31], where $\tau$ runs from $-J,-J+1, \ldots, J-1,+J$, with energy increasing from $\tau=-J$. The $+/-$ parity of an asymmetric top molecule such as $\mathrm{H}_{2}^{32} \mathrm{~S}$ is defined by $(-1)^{K_{c}}$ [19]. $\mathrm{H}_{2}^{32} \mathrm{~S}$ belongs to the $C_{2 \mathrm{v}}(\mathrm{M})$ molecular symmetry (MS) group [32], which contains irreducible representations $\mathrm{A}_{1}, \mathrm{~A}_{2}, \mathrm{~B}_{1}$, and $\mathrm{B}_{2}$, as given in Table 2.

### 2.3. Selection rules

The rigorous selection rules governing rotation-vibration transitions for a molecule of the $C_{2 v}(\mathrm{M}) \mathrm{MS}$ group are given by:
$\Delta J=0, \pm 1$,
$J^{\prime}+J^{\prime \prime} \neq 0$,
$-\leftrightarrow+$

Table 1
Quantum numbers used to label the upper and lower energy states of $\mathrm{H}_{2}^{32} \mathrm{~S}$.

| Label | Description |
| :--- | :--- |
| $v_{1}$ | S-H symmetric stretch $\left(\sim 2614.4 \mathrm{~cm}^{-1}\right)$ |
| $v_{2}$ | Symmetric bending mode $\left(\sim 1182.6 \mathrm{~cm}^{-1}\right)$ |
| $v_{3}$ | S-H antisymmetric stretch $\left(\sim 2628.5 \mathrm{~cm}^{-1}\right)$ |
| $J$ | Rotational angular momentum |
| $K_{a}, K_{c}$ | Projections of rotational angular momentum |
| ortho/para | Nuclear spin state (see text) |

Table 2
Symmetry of the rovibrational states of $\mathrm{H}_{2}^{32} \mathrm{~S}$.

| Symmetry | $\mathrm{A}_{1}$ | $\mathrm{~A}_{2}$ | $\mathrm{~B}_{1}$ | $\mathrm{~B}_{2}$ |
| :--- | :--- | :--- | :--- | :--- |
| Parity | + | - | - | + |
| Nuclear spin state | para | para | ortho | ortho |

The ortho states of $\mathrm{H}_{2}^{32}$ S have the nuclear spin statistical weight $g_{\mathrm{ns}}=3$, while for the para states $g_{\mathrm{ns}}=1$, thus, ortho transitions have three times the intensity of para transitions. This is sometimes referred to as intensity alternation. It is assumed that ortho and para states do not interconvert. Such transitions are very weakly allowed [33] but have yet to be observed for $\mathrm{H}_{2}^{32} \mathrm{~S}$.

## 3. Experimental sources

A large number of experimentally determined transition wavenumbers can be found in the literature for the main isotopologue of hydrogen sulfide, $\mathrm{H}_{2}^{32} \mathrm{~S}$. We have attempted to conduct a rigorous and comprehensive search for all useable spectroscopic data. Fortunately, much of the data up to 2012 was previously collated as part of the 2012 W@DIS information system for hydrogen sulfide [14], in which some of the authors of this paper were involved. These data were converted to Marvel format for this work and analysed alongside data from newly collected sources. This requires the transition wavenumber (in $\mathrm{cm}^{-1}$ ) and the associated uncertainty, along with quantum number assignments for both the upper and lower energy states, and a unique reference label for each transition. This reference indicates the data source the transition originates from. The data source tag is based on the notation employed by an IUPAC Task Group on water spectroscopy [19]. An extract of the input file in the required format is given in Table 3; the full file can be found in the supplementary data for this publication.

33 sources of experimental data were used in the final data set. The data from more recent papers are generally provided in digital format, but some of the older papers had to be processed through digitalisation software, or even manually entered in the worst cases. After digitalisation the data were converted to Marvel format, as described above.

Table 4 gives a summary of all the data sources used in this work, along with the energy range, number of transitions (as-

Table 3
Extract from the Marvel input file for the ortho transitions for $\mathrm{H}_{2}^{32} \mathrm{~S}$. The full file is supplied as part of the supplementary information to this paper. All energy term values and uncertainties are in units of $\mathrm{cm}^{-1}$. The meaning of the upper and lower state assignments can be found in Table 1.

| Transition | Uncertainty | Upper state assignment | Lower state assignment | Reference |
| :--- | :--- | :--- | :--- | :--- |
| 33.12631 | 0.00015 | 000523 ortho | 000514 ortho | 94YaKl_1 |
| 33.12631 | 0.00018 | 000303 ortho | 000212 ortho | 94YaKl_2 |
| 34.00529 | 0.00010 | 000872 ortho | 000863 ortho | 94YaKl_5 |
| 34.15779 | 0.00010 | 0001192 ortho | 0001183 ortho | 94YaKl_7 |
| 34.21980 | 0.00010 | 000643 ortho | 000634 ortho | 94YaKl_8 |
| 34.24062 | 0.00012 | 000423 ortho | 000414 ortho | 94YaKl_9 |
| 35.73512 | 0.00010 | 000863 ortho | 000854 ortho | 94YaKl_13 |

Table 4
Data sources used in this study with frequency range, numbers of transitions ( $\mathrm{A} / \mathrm{V}$ for assigned/verified), and comments, which are detailed in Section 3.1. As far as we are aware all experiments were conducted at room temperature.

| Tag | Ref. | Range $\left(\mathrm{cm}^{-1}\right)$ | A/V | Comments |
| :--- | :--- | :--- | :--- | :--- |
| 72HeCoLu | $[34]$ | $1.17-25.55$ | $37 / 35$ | $(4 \mathrm{a})$ |
| 95BeYaWiPo | $[35]$ | $4.39-85.41$ | $112 / 84$ | $(4 \mathrm{~b})$ |
| 68CuKeGa | $[36]$ | $5.63-14.15$ | $6 / 6$ |  |
| 71Huiszoon | $[37]$ | $5.63-7.23$ | $2 / 2$ |  |
| 14CaPu | $[38]$ | $7.23-53.17$ | $70 / 70$ |  |
| 85BuFeMeSh | $[39]$ | $10.02-20.9$ | $6 / 6$ |  |
| 94YaKl | $[40]$ | $33.13-259.76$ | $366 / 366$ |  |
| 13CaPu | $[41]$ | $33.97-37.45$ | $4 / 4$ |  |
| 13AzYuTeMa | $[42]$ | $45.25-359.79$ | $1158 / 1139$ | $(4 \mathrm{c})$ |
| 83FlCaJo | $[43]$ | $50.77-307.51$ | $426 / 387$ | $(4 \mathrm{~d})$ |
| 18UlBeGr | $[44]$ | $729.78-1735.41$ | $2267 / 2267$ |  |
| 82LaEdGiBo | $[45]$ | $1003.46-1495.28$ | $397 / 396$ | $(4 \mathrm{e})$ |
| 83Strow | $[46]$ | $1082.03-1257.07$ | $123 / 123$ |  |
| 96UlMaKoAl | $[47]$ | $1178.05-1359.78$ | $41 / 41$ |  |
| 98BrCrCrNa | $[48]$ | $2141.30-4249.85$ | $7473 / 7473$ | $(4 \mathrm{f}$ |
| 18Horneman | $[49]$ | $2180.35-4220.46$ | $4460 / 4460$ |  |
| 84LeFlCaJo | $[50]$ | $2180.36-2945.81$ | $2113 / 2111$ | $(4 \mathrm{f})$ |
| 81GiEd | $[51]$ | $2192.48-2823.11$ | $715 / 704$ | $(4 \mathrm{~g})$ |
| 96UlOnKoAl | $[52]$ | $3614.70-3887.66$ | $106 / 106$ |  |
| 05UlLiBeGr | $[53]$ | $4000.59-6653.79$ | $2347 / 2347$ |  |
| 97BrCrCrNa | $[54]$ | $4500.88-5595.01$ | $5221 / 5219$ | $(4 \mathrm{~h})$ |
| 18Liu | $[55]$ | $4514.79-5555.58$ | $3337 / 3335$ | $(4 \mathrm{i})$ |
| 04BrNaPoSi_c | $[56]$ | $5688.27-6676.71$ | $3178 / 3178$ |  |
| 04BrNaPoSi_a | $[57]$ | $7169.19-7898.97$ | $2878 / 2876$ | $(4 \mathrm{j})$ |
| 04UlLiBeGr_b | $[58]$ | $7226.81-7994.09$ | $1855 / 1855$ |  |
| 04UlLiBeGr_a | $[59]$ | $8405.91-8905.28$ | $589 / 589$ |  |
| 04BrNaPoSi_b | $[60]$ | $8412.73-8906.11$ | $1179 / 1175$ | $(4 \mathrm{k})$ |
| 03DiNaHuZh | $[61]$ | $9541.01-10000.71$ | $1736 / 1728$ | $(4 \mathrm{l})$ |
| 01NaCa_a | $[62]$ | $10787.33-11297.99$ | $1105 / 1097$ | $(4 \mathrm{~m})$ |
| 94GrRaStDe | $[63]$ | $11948.91-12246.28$ | $227 / 145$ | $(4 \mathrm{n})$ |
| 97VaBiCaFl | $[64]$ | $12324.55-12670.68$ | $399 / 387$ | $(4 \mathrm{o})$ |
| 99CaFl | $[65]$ | $13060.51-13357.14$ | $219 / 206$ | $(4 \mathrm{p})$ |
| 01NaCa_b | $[66]$ | $16186.25-16436.57$ | $173 / 154$ | $(4 \mathrm{q})$ |
| Total |  | $1.17-16436.57$ | $44325 / 44071$ |  |
|  |  |  |  |  |

signed (A) and verified (V)), and comments, which are detailed in Section 3.1. Table 5 lists those data sources which were considered but not used, with comments on the reasons for their exclusion from the analysis. The reference tag given in these tables matches those used in the unique labels in the Marvel input files, given in the supplementary data and illustrated in Table 3.

As transitions have never been observed between ortho and para states of $\mathrm{H}_{2}^{32} \mathrm{~S}$, they form two separate principal components (PCs) of the experimental spectroscopic network. All input and out-
put files supplied in the supplementary data to this work are split into either ortho or para.

### 3.1. Comments on the experimental sources of Table 4

(4a) 72 HeCoLu [34] contains 2 lines which have been cited as taken from other experimental sources, for which the original data are already in our dataset. These duplicates were removed.
(4b) 95 BeYaWiPo [35] contains 28 lines which have been cited as from other experimental sources, for which the original data are already in our dataset. These duplicates were removed.
(4c) 13AzYuTeMa [42] gives data for rotational lines recorded at room temperature. The equipment used provides spectra with very high sensitivity and this enabled transitions between highly rotationally excited states to be recorded, even at room temperature. These high rotational excitations may not be so accurately assigned as those of lower energy, as there are no data from other sources in the same region to confirm these measurements. The supplementary data from the original paper does not contain the original experimental transitions; this was provided by the corresponding author. There were 19 lines which could not be validated against other data and so were removed from our dataset (see Section 3.2).
(4d) 83FlCaJo [43] contains 39 lines which have been cited as from other sources which are already present in our dataset. The duplicates were removed.
(4e) 82LaEdGiBo [45] contains one line which could not be validated and so has been removed from our dataset.
(4f) 84LeFlCaJo [50] contains 2 lines which could not be validated against other more recent data and were removed from our dataset. All data from this source were recalibrated; the calibration factor employed was 0.999999746 below $2450 \mathrm{~cm}^{-1}$ and 1.0000000714 above $2450 \mathrm{~cm}^{-1}$.
( $\mathbf{4 g}$ ) 81 GiEd [51] contains 11 lines which could not be validated against more recent data and thus were removed from our dataset.
(4h) 97 BrCrCrNa [54] contains 2 lines with no corresponding energy in the AYT2 linelist (see Section 5) and so were removed from our dataset.
(4i) 18Liu [55] contains 2 lines with no corresponding energy in the AYT2 linelist and so were removed from our dataset.
(4j) 04BrNaPoSi_a [57] contains 2 lines with no corresponding energy in the AYT2 linelist and so were removed from our dataset.
(4k) 04 BrNaPoSi b [60] contains 4 lines which could not be validated and were removed from our dataset.
(41) 03DiNaHuZh [61] contains one blended line which has been removed from our dataset. A further 7 lines were found to include

Table 5
Data sources considered but not used in this work.

| Tag | Ref. | Comments |
| :--- | :--- | :--- |
| 02CoRoTy | $[67]$ | Data taken from other sources |
| 94WaKuSu | $[68]$ | Data taken from other sources |
| 96SuMeKr | $[69]$ | Data taken from other sources |
| 97SuMeKr | $[70]$ | Data taken from other sources |
| 97Sumpf | $[71]$ | Data taken from other sources |
| 98PiPoCoDe | $[72]$ | Data from http://spec.jpl.nasa.gov. Data does not appear to be experimental. |
| 73HeDeKi | $[73]$ | Data taken from other sources, [34,36,37] |
| 02KiSuKrTi | $[74]$ | Data appears to be taken from the 1996 edition of HITRAN [75] |
| 06Polovtseva | $\mathrm{N} / \mathrm{A}$ | Thesis, no citation. Referenced in [14] |
| 87LeFlCaAr | $[76]$ | Energy levels |
| 13Azzam | $[77]$ | All results are given in 13AzYuTeMa [42] |
| 94KoJe | $[78]$ | Energy levels |
| 95FlGrRa | $[79]$ | Energy levels |
| 85LaEdGiBo | $[80]$ | Calculated values, compared to experimental values from 83FlCajo [43] |
| 94ByNaSmSi | $[81]$ | Energy Levels |
| 01TyTaSc | $[82]$ | Vibrational energy Levels |
| 69MiLeHa | $[83]$ | Could not be validated with the other sources to a reasonable accuracy (see Section 3.2) |
| 56AIPl | $[31]$ | Could not be validated with the other sources to a reasonable accuracy (see Section 3.2) |
| 69SnEd | $[84]$ | Could not be validated with the other sources to a reasonable accuracy (see Section 3.2) |

a level with no corresponding energy in the AYT2 linelist and so were removed from our dataset.
( $\mathbf{4 m}$ ) 01 NaCa a [62] contains 8 transitions which were found to have no corresponding levels in the AYT2 linelist and so were removed from our dataset.
(4n) 94GrRaStDe [63] tentatively assigned the vibrational band in their data as $\left(v_{1}, v_{2}, v_{3}\right)=(2,2,2)$; however, in 95FIGrRaSt [79], a paper published later with the same authors, the revised assignment is $\left(v_{1}, v_{2}, v_{3}\right)=(3,0,2)$. We adopt the latter vibrational label in our dataset. No other data sources probe this vibrational band. The frequency is only given to 3 decimal places, so the uncertainty was altered to match. The original dataset has 82 transitions either labelled with an asterisk to indicate lines from the $\mathrm{H}_{2}^{34} \mathrm{~S}$ isotopologue, or labelled with a dagger to indicate lines which have been perturbed due to those from $\mathrm{H}_{2}^{34} \mathrm{~S}$ [63]. We have commented these out by adding a minus sign to the wavenumber and added the label $\_p t$ to the end.
(40) 97VaBiCaFl [64] contains 4 lines which could not be validated and so were removed from our dataset. A further 8 lines were found to contain a level with no corresponding energy in the AYT2 linelist and so were removed from our dataset.
(4p) The assignment for 99CaFl_185_na_ct from 99CaFl [65] leads to different spin states (ortho/para) for the upper and lower state, which is forbidden. We have removed this transition from our dataset. 5 other lines could not be validated and so were removed from our dataset. 7 lines were found to have no variational counterparts in the AYT2 linelist and so were removed.
(4q) 01 NaCa b [66] contains one line which could not be validated and so was removed from our dataset. A further 18 lines were found to contain levels with no corresponding energies in the AYT2 linelist and so were removed from our dataset.

### 3.2. General comments

All transitions which were considered but not processed in the final dataset have a minus sign in front of the transition wavenumber (indicating that Marvel will ignore them) in the input files provided as supplementary material and are labelled with a comment at the end of the reference (see the comments of Section 3.1). We used a cut-off of $0.035 \mathrm{~cm}^{-1}$ as the largest acceptable uncertainty and removed any transitions with an uncertainty greater than this, with a note in Section 3.1 to indicate how many transitions from a particular data source could not be validated to within this accuracy.

The publications 97 BrCrCrNa [54], 04BrNaPoSi_c [56], 04BrNaPoSi_a [57], and 04BrNaPoSi_b [60] include just a short description of the assignment and modelling of the Fourier Transform (FT) spectra between 4500 and $8900 \mathrm{~cm}^{-1}$. Detailed information on the theoretical treatment of these spectra will be published separately in due course. Some of these transitions, between 4400$8000 \mathrm{~cm}^{-1}$, were previously reported in the HITRAN-2012 [85] and HITRAN-2016 [86] databases, while the second decade region, $8400-8900 \mathrm{~cm}^{-1}$, is presented for the first time. It is also worth mentioning that the Marvel dataset includes a considerable number of transitions which include new, unpublished energy levels. As only the upper energy levels of the transitions were reported in 05UlLiBeGr [53], 04UlLiBeGr_b [58], and 04UlLiBeGr_a [59], the corresponding transition wavenumbers were recovered from Liu (18Liu [55]) using the energy levels obtained from 97 BrCr CrNa [54], 04BrNaPoSi_c [56], 04BrNaPoSi_a [57], and 04BrNaPoSi_b [60]. For this reason, the transitions in the Marvel dataset which are referred to as from 05UlLiBeGr [53], 04UlLiBeGr_b [58], and 04UlLiBeGr_a [59] contain a larger number of assignments than reported in the original publications. The data of the first hexad region between $4500-5600 \mathrm{~cm}^{-1}$ were totally assigned based on the energy levels reported in 97 BrCrCrNa [54]. The highly
accurate (an accuracy of $0.0005 \mathrm{~cm}^{-1}$ ) set of $\mathrm{H}_{2}^{32} \mathrm{~S}$ transitions recorded using a FT spectrometer between $2200-4250 \mathrm{~cm}^{-1}$ was provided by Horneman (18Horneman [49]) and assigned using the energy levels reported in 98 BrCrCrNa [48].

### 3.3. Sources from the HITRAN database

$\mathrm{H}_{2}^{32} \mathrm{~S}$ has been included in the HITRAN database [85-90] since 1991 [87]. The following are sources for the line positions and energy levels of the $H_{2}^{32} \mathrm{~S}$ data in the HITRAN database, up to the 2016 release: 96UlMaKoAl [47], 83FICaJo [43], 98 BrCrCrNa [48], 13AzYuTeMa [42], 94YaKl [40], 95BeYaWiPo [35], 05UlLiBeGr [53], 04UlLiBeGr [58], 03DiNaHuZh [61], 01NaCa_b [66], 84LeFICajo [50], 82LaEdGiBo [45], 85LaEdGiBo [80], 94ByNaSmSi [81]. The variational ExoMol AYT2 linelist [91] was used to assign transitions in the $v_{2}$ excited vibrational state in the 2012 release [85].

## 4. Results

The Marvel website (http://www.kkrk.chem.elte.hu/ marvelonline) has a version of Marvel which can be run online. The variable NQN (number of quantum numbers) is 7 in the case of hydrogen sulfide, as illustrated in Table 3 which shows an extract of the ortho input file to Marvel.

Marvel automatically assigns the lowest energy state in a particular component of the spectroscopic network to 0 . The ground rovibrational state is included in the para set of energy levels, however there needs to be a "magic number", corresponding to the energy of the lowest ortho state, which is added to all the Marvel ortho-symmetry energies. Here, this was taken as the ground vibrational state $\left(v_{1}, v_{2}, v_{3}\right)=(0,0,0)$ with the lowest rotational energy (see Section 2.2), $J=1, K_{a}=0, K_{c}=1$, of $94 K o J e$ [78], who determined the value of $13.74631 \mathrm{~cm}^{-1}$. The output for the ortho energies in the supplementary data, and the extract in Table 6 , all have this magic number added for the ortho principal spectroscopic network. There are a small number (4 para) of energy levels which are not joined to either of the principal components. If more experimental transitions became available in the future it would be possible to link these to the para principal network.

We collated and considered a total of 44325 transitions from 33 experimental sources ( 25474 ortho and 18851 para). Of those 254 were found to be inconsistent (could not be validated to within $0.035 \mathrm{~cm}^{-1}$ ) with others and so removed from the final data set, leaving a total of 44071 transitions used as input into Marvel (25 293 ortho and 18778 para).

Fig. 1 gives a visual representation of NumTrans, the number of transitions which link the states of the Marvel ortho $-\mathrm{H}_{2}^{32} \mathrm{~S}$ network, against corresponding upper state energies. Those upper states which are dark blue in colour, linked by only 1 transition, should be considered less reliable than those in red, which are supported by hundreds of different transitions, up to 404 . The values of NumTrans for each level are given in the energy level files in the supplementary data.

## 5. Comparison to other derived energy levels

We used a number of sources to compare energy level values against those determined in this work. 94KoJe [78] contains a list of energy levels, which are taken or calculated from other sources [31,43,50,76,80,84]; see Table 7 for a breakdown by vibrational band. AYT2 [91] is a theoretical variational linelist for $\mathrm{H}_{2}{ }^{32} \mathrm{~S}$ calculated as part of the ExoMol project [92,93], a database of theoretical linelists for molecules of astrophysical importance. AYT2 is appropriate up to temperatures of around 2000 K and designed for use in characterising the atmospheres of cool stars and exoplanets. The states file, available from https://www.exomol.com,

Table 6
Extract from the Marvel output file for the ortho energy levels of $\mathrm{H}_{2}^{32}$ S. The full file is supplied as part of the supplementary information to this paper. All energies and uncertainties are in units of $\mathrm{cm}^{-1}$. NumTrans gives the number of transitions which are linked to that particular energy level. Assignments are as given by Table 1.

| Assignment |  |  |  |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  | Energy | Uncertainty | NumTrans | Sym |  |  |  |  |
| 0 | 0 | 0 | 1 | 0 | 1 | ortho | 13.746310 | 0.000001 | 177 | B1 |
| 0 | 0 | 0 | 1 | 1 | 0 | ortho | 19.375630 | 0.000001 | 153 | B2 |
| 0 | 0 | 0 | 2 | 1 | 2 | ortho | 38.297765 | 0.000002 | 271 | B2 |
| 0 | 0 | 0 | 2 | 2 | 1 | ortho | 55.161605 | 0.000002 | 265 | B1 |
| 0 | 0 | 0 | 3 | 0 | 3 | ortho | 71.424233 | 0.000002 | 282 | B1 |
| 0 | 0 | 0 | 3 | 1 | 2 | ortho | 95.056325 | 0.000001 | 344 | B2 |
| 0 | 0 | 0 | 3 | 2 | 1 | ortho | 107.368228 | 0.000001 | 331 | B1 |
| 0 | 0 | 0 | 4 | 1 | 4 | ortho | 114.177613 | 0.000001 | 282 | B2 |
| 0 | 0 | 0 | 3 | 3 | 0 | ortho | 117.392015 | 0.000001 | 265 | B2 |
| 0 | 0 | 0 | 4 | 2 | 3 | ortho | 148.418340 | 0.000001 | 379 | B1 |



Fig. 1. The lower-state energies of the experimental ortho $-\mathrm{H}_{2}^{32} \mathrm{~S}$ transitions used in this work, against corresponding upper-state energies. The vertical bars along the horizontal-axis show the lower-state energies, while the horizontal bars along the vertical-axis give the upper-state energies. Each circle represents a particular transition, with the size proportional to the log of NumTrans, the number of transitions supporting the upper state. This value ranges from 1 (dark blue) to 404 (red). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
gives the calculated energy levels, which can be compared against our empirical energy levels. For more information on the format of the ExoMol data files, please refer to [93]. 95FlGrRaSt [79] contains rovibrational energy levels from the $\left(v_{1}, v_{2}, v_{3}\right)=(2,0,3)$ and $(3,0,2)$ vibrational bands. The authors outline an experimental procedure used to deduce these levels but they do not provide the experimental transition data so we could not include the data from this source in our dataset. 87LeFlCaAr [76] has a table of rovibrational energy levels for the $\left(v_{1}, v_{2}, v_{3}\right)=(2,1,0),(1,1,1)$ and ( $0,1,2$ ) vibrational bands. 96UlMaKoAl [47] contains rovibrational energy levels from the $\left(v_{1}, v_{2}, v_{3}\right)=(0,1,0)$ vibrational band.

We compared the pure rotational levels determined in this work both against those given in 94 KoJe [78] and the calculated values from the AYT2 linelist [91], up to $J=5$. These data are given in Table 8. The pure rotational levels are closer to those from the experimentally determined sources of 94KoJe than to those of variationally calculated AYT2, as would be expected.

The pure vibrational levels from this work are also compared with those from 94KoJe [78] and AYT2 [91], see Table 9. It should be noted here that there are some differences in labelling between these data sources and those used in this work. For example, the pure vibrational levels of 94 KoJe [78] which are labelled $\left(v_{1}, v_{2}, v_{3}\right)=(2,0,2),(1,0,2)$ and ( $3,0,0$ ), match the normal mode labelling of ( $4,0,0$ ), ( $3,0,0$ ) and ( $1,0,2$ ), respectively,


Fig. 2. Deviations in $\mathrm{cm}^{-1}$ between energy levels from this work and: 94Koje [78] (see Table 7), 95FlGrRaSt [79], 87LeFlCaAr [76], 96UlMaKoAl [47]. Note the logarithmic vertical-axis.
according to the labelling of 01TyTaSc [82] and the experimental sources used in this work.

We compared all the rovibrational energy levels given in 94KoJe [78], 95FlGrRaSt [79], 96UlMaKoAl [47], and 87LeFlCaAr [76] against our set of Marvel energy levels, taking the aforementioned labelling differences into account for 94KoJe [78] (see Table 7). These comparisons are illustrated in Fig. 2. Most levels are in good agreement, with only a few levels with differences between $0.1-0.4 \mathrm{~cm}^{-1}$. These levels with the largest differences are from 87LeFlCaAr [76] and 56AIPl [31]. All the levels which are compared to those of 87 LeFlCaAr [76] have several transitions which link them to other energy levels in our Marvel dataset, and originate from different sources, which increases their reliability. This indicates that the lack of accuracy for these particular energies originates from 87LeFlCaAr [76] and not the Marvel dataset. The transition data from 56AlPl [31] was considered for use in the current study, but the data could not be validated with that from newer sources (see Table 5), and so the energy levels are also assumed to be unreliable.

As mentioned above, AYT2 [91] is a theoretical variational linelist for $\mathrm{H}_{2}^{32} \mathrm{~S}$ calculated as part of the ExoMol project [92,93]. Highly accurate experimental energy levels provide essential input for testing and improving theoretically calculated line positions such as those within AYT2. The ExoMol data format convention is to have a 'states' and a 'trans' file. The states file gives the calculated rovibrational energy levels, which can be compared against our empirical energy levels, and the trans file contains the Einstein-A coefficients for transitions between these states [93].

Table 7
Original sources of energy level data from 94KoJe [78].

| Tag | Ref. | Vibrational bands | Comments |
| :---: | :---: | :---: | :---: |
| 83FlCajo | [43] | $(0,0,0)$ | Experimental source used in this work |
| 85LaEdGiBo | [80] | $(0,1,0)$ | Calculated values, compared to experimental values from 83FICaJo [43] |
| 84LeFlCajo | [50] | $(0,2,0)(1,0,0)(0,0,1)$ | Experimental source used in this work |
| 69SnEd | [84] | $(1,1,0)(0,1,1)$ | Experimental source used in this work |
| 56AlPl | [31] | $\begin{aligned} & (0,2,1)(2,0,0)(1,0,1)(3,0,0)(2,0,1)(1,0,2)(0,0,3)(2,1,1) \\ & \quad(3,0,1)(2,0,2)(1,0,3)(3,1,1) \end{aligned}$ | Data source not considered reliable. All $J=0$ levels only. |
| 87LeFlCaAr | [76] | $(2,1,0)(1,1,1)$ | 94KoJe [78] only contains levels from this source up to $J=5$. See text and Fig. 2 for a full comparison. |

Table 8
Comparison of pure rotational levels from this work with those of 94KoJe [78] (see Table 7) and AYT2 [91] up to $J=5$. NumTrans gives the number of transitions linking a particular state. All energies and uncertainties are in $\mathrm{cm}^{-1}$. The uncertainty refers to this work.

| $J K_{a} K_{c}$ | State | Sym | NumTrans | Uncertainty | This work | 94KoJe | Difference | AYT2 | Difference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 000 | para | $\mathrm{A}_{1}$ | 58 | 0.000098 | 0 | 0 | 0 | 0 | 0 |
| 101 | ortho | $\mathrm{B}_{1}$ | 170 | 0.000001 | 13.746310 | 13.74631 | 0.00000 | 13.746340 | 0.000030 |
| 111 | para | $\mathrm{A}_{2}$ | 157 | 0.000002 | 15.090119 | 15.09011 | -0.00001 | 15.090151 | 0.000032 |
| 110 | ortho | $\mathrm{B}_{2}$ | 147 | 0.000001 | 19.375630 | 19.37563 | 0.00000 | 19.375563 | -0.000067 |
| 202 | para | $\mathrm{A}_{1}$ | 223 | 0.000001 | 38.016095 | 38.01607 | -0.00003 | 38.016264 | 0.000169 |
| 212 | ortho | $\mathrm{B}_{2}$ | 258 | 0.000002 | 38.297765 | 38.29775 | -0.00001 | 38.297961 | 0.000196 |
| 211 | para | $\mathrm{A}_{2}$ | 211 | 0.000001 | 51.140188 | 51.14016 | -0.00003 | 51.140060 | -0.000128 |
| 221 | ortho | $\mathrm{B}_{1}$ | 253 | 0.000002 | 55.161605 | 55.16158 | -0.00003 | 55.161460 | -0.000145 |
| 220 | para | $\mathrm{A}_{1}$ | 181 | 0.000001 | 58.368870 | 58.36884 | -0.00003 | 58.368621 | -0.000249 |
| 303 | ortho | $\mathrm{B}_{1}$ | 265 | 0.000002 | 71.424221 | 71.42426 | 0.00004 | 71.424760 | 0.000539 |
| 313 | para | $\mathrm{A}_{2}$ | 213 | 0.000001 | 71.465192 | 71.46515 | -0.00004 | 71.465653 | 0.000461 |
| 312 | ortho | $\mathrm{B}_{2}$ | 329 | 0.000001 | 95.056313 | 95.05630 | -0.00001 | 95.056267 | -0.000046 |
| 322 | para | $\mathrm{A}_{1}$ | 272 | 0.000001 | 96.392529 | 96.39246 | -0.00007 | 96.392428 | -0.000101 |
| 321 | ortho | $\mathrm{B}_{1}$ | 313 | 0.000001 | 107.368216 | 107.36820 | -0.00002 | 107.367927 | -0.000289 |
| 404 | para | $\mathrm{A}_{1}$ | 205 | 0.000001 | 114.172246 | 114.17217 | -0.00008 | 114.173085 | 0.000839 |
| 414 | ortho | $\mathrm{B}_{2}$ | 270 | 0.000001 | 114.177601 | 114.17758 | -0.00002 | 114.178495 | 0.000894 |
| 331 | para | $\mathrm{A}_{2}$ | 250 | 0.000002 | 115.340656 | 115.34059 | -0.00007 | 115.340038 | -0.000618 |
| 330 | ortho | $\mathrm{B}_{2}$ | 247 | 0.000001 | 117.392003 | 117.39199 | -0.00001 | 117.391407 | -0.000596 |
| 413 | para | $\mathrm{A}_{2}$ | 306 | 0.000001 | 148.140653 | 148.14058 | -0.00007 | 148.140740 | 0.000087 |
| 423 | ortho | $\mathrm{B}_{1}$ | 362 | 0.000001 | 148.418328 | 148.41831 | -0.00002 | 148.418470 | 0.000142 |
| 505 | ortho | $\mathrm{B}_{1}$ | 264 | 0.000045 | 166.343488 | 166.34345 | -0.00004 | 166.344887 | 0.001399 |
| 515 | para | $\mathrm{A}_{2}$ | 196 | 0.000077 | 166.344005 | 166.34417 | 0.00016 | 166.345606 | 0.001601 |
| 422 | para | $\mathrm{A}_{1}$ | 308 | 0.000001 | 170.335799 | 170.33574 | -0.00006 | 170.335471 | -0.000328 |
| 432 | ortho | $\mathrm{B}_{2}$ | 383 | 0.000002 | 173.967266 | 173.96726 | -0.00001 | 173.966794 | -0.000472 |
| 431 | para | $\mathrm{A}_{2}$ | 261 | 0.000001 | 182.648548 | 182.64849 | -0.00006 | 182.647978 | -0.000570 |
| 441 | ortho | $\mathrm{B}_{1}$ | 290 | 0.000002 | 195.661414 | 195.66142 | 0.00001 | 195.659965 | -0.001449 |
| 440 | para | $\mathrm{A}_{1}$ | 219 | 0.000003 | 196.802189 | 196.80212 | -0.00007 | 196.800713 | -0.001476 |
| 514 | ortho | $\mathrm{B}_{2}$ | 356 | 0.000037 | 210.217262 | 210.21727 | 0.00001 | 210.217731 | 0.000469 |
| 524 | para | $\mathrm{A}_{1}$ | 270 | 0.000001 | 210.264821 | 210.26477 | -0.00005 | 210.265229 | 0.000408 |
| 523 | ortho | $\mathrm{B}_{1}$ | 387 | 0.000001 | 243.343434 | 243.34344 | 0.00001 | 243.343211 | -0.000223 |
| 533 | para | $\mathrm{A}_{2}$ | 333 | 0.000001 | 244.392517 | 244.39250 | -0.00002 | 244.392149 | -0.000368 |
| 532 | ortho | $\mathrm{B}_{2}$ | 363 | 0.000001 | 263.738931 | 263.73895 | 0.00002 | 263.738560 | -0.000371 |
| 542 | para | $\mathrm{A}_{1}$ | 298 | 0.000001 | 271.106067 | 271.10604 | -0.00003 | 271.104809 | -0.001258 |
| 541 | ortho | $\mathrm{B}_{1}$ | 316 | 0.000002 | 277.337562 | 277.33758 | 0.00002 | 277.336633 | -0.000929 |
| 551 | para | $\mathrm{A}_{2}$ | 229 | 0.000002 | 296.104442 | 296.10442 | -0.00002 | 296.101282 | -0.003160 |
| 550 | ortho | $\mathrm{B}_{2}$ | 278 | 0.000001 | 296.677577 | 296.67760 | 0.00002 | 296.674560 | -0.003017 |

This format allows calculated linelists to be retrospectively updated using more reliable experimental energy levels in order to improve their accuracy, see [94] for an example.

To compare our energy levels with those from the AYT2 [91] linelist, a comparable states file was made; we labelled our states with the same $A_{1}, A_{2}, B_{1}, B_{2}$ symmetry labels which are given in the AYT2 states file, see Table 2. We then matched all states with this same symmetry, $v_{2}, v_{1}+v_{3}$, and $J$, and searched for the closest value within these given parameters. Fig. 3 gives the result of this comparison.

Work is underway to update the existing ExoMol AYT2 states file and linelist based on the energy levels obtained in this work, using only those levels based on the transitions we are most sure about. Care should be taken in general when using Marvel energy levels with a low value of NumTrans (see Fig. 1). This, along with the uncertainty, gives an indication of the reliability of a particular energy level.

## 6. Conclusions

A total of 44325 measured experimental rovibrational transitions of $\mathrm{H}_{2}^{32} \mathrm{~S}$ from 33 publications have been considered in this work. From this set, 3969 ortho- and 3467 para- $\mathrm{H}_{2}^{32}$ S energy levels have been determined using the Measured Active RotationalVibrational Energy Levels (Marvel) technique. These results have been carefully compared with alternative compilations of energy levels.

A variational high-temperature linelist for $\mathrm{H}_{2}^{32} \mathrm{~S}$ has been computed as part of the ExoMol project, AYT2 [91]. Our new Marvel energy levels will be used to improve the accuracy of this theoretical linelist.

A significant part of this work was performed by pupils from Highams Park School in London, as part of a project known as ORBYTS (Original Research By Young Twinkle Students) [95]. The Marvel studies of ${ }^{12} \mathrm{C}_{2} \mathrm{H}_{2}$ [26] and ${ }^{48} \mathrm{Ti}^{16} \mathrm{O}$ [18] were also per-

Table 9
Comparison of pure vibrational levels $(J=0)$ from this work with those of 94KoJe [78] (see Table 7) and AYT2 [91]. NT stands for NumTrans, the number of transitions linking a particular state. All energies and uncertainties are in $\mathrm{cm}^{-1}$. The uncertainty refers to this work. State is from this work and Sym is the corresponding symmetry label as used in AYT2 [91] (see Section 2.2).

| $\nu_{1} \nu_{2} \nu_{3}$ | State | Sym | NT | Uncertainty | This work | 94KoJe | Difference | AYT2 | Difference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 000 | para | $\mathrm{A}_{1}$ | 58 | 0.000098 | 0 |  |  | 0 | 0 |
| 010 | para | $\mathrm{A}_{1}$ | 5 | 0.000477 | 1182.576991 | 1182.5742 | -0.0028 | 1182.569618 | -0.007373 |
| 020 | para | $\mathrm{A}_{1}$ | 4 | 0.000629 | 2353.964679 | 2353.9655 | 0.0008 | 2353.907317 | -0.057362 |
| 100 | para | $\mathrm{A}_{1}$ | 3 | 0.000663 | 2614.407743 | 2614.4074 | -0.0003 | 2614.394829 | -0.012914 |
| 001 | ortho | $\mathrm{B}_{2}$ | 2 | 0.000669 | 2628.454821 | 2628.4552 | 0.0004 | 2628.463320 | 0.008499 |
| 030 | para | $\mathrm{A}_{1}$ | 3 | 0.000621 | 3513.789974 |  |  | 3513.705072 | -0.084902 |
| 110 | para | $\mathrm{A}_{1}$ | 3 | 0.000390 | 3779.166566 | 3779.1710 | 0.0044 | 3779.189348 | 0.022782 |
| 011 | ortho | $\mathrm{B}_{2}$ | 2 | 0.000625 | 3789.269211 | 3789.2720 | 0.0028 | 3789.269878 | 0.000667 |
| 040 | para | $\mathrm{A}_{1}$ | 1 | 0.001000 | 4661.672219 |  |  | 4661.605794 | -0.066425 |
| 120 | para | $\mathrm{A}_{1}$ | 2 | 0.000707 | 4932.699369 |  |  | 4932.688937 | -0.010432 |
| 021 | ortho | $\mathrm{B}_{2}$ | 2 | 0.000707 | 4939.104010 | 4939.2300 | 0.1260 | 4939.129851 | 0.025841 |
| 200 | para | $\mathrm{A}_{1}$ | 2 | 0.000707 | 5144.986319 | 5145.1200 | 0.1337 | 5145.031868 | 0.045549 |
| 101 | ortho | $\mathrm{B}_{2}$ | 2 | 0.000707 | 5147.220560 | 5147.3600 | 0.1394 | 5147.166622 | -0.053938 |
| 002 | para | $\mathrm{A}_{1}$ | 2 | 0.000707 | 5243.101919 |  |  | 5243.158956 | 0.057037 |
| 130 | para | $\mathrm{A}_{1}$ | 2 | 0.000928 | 6074.581067 |  |  | 6074.566059 | -0.015008 |
| 031 | ortho | $\mathrm{B}_{2}$ | 2 | 0.000707 | 6077.594560 |  |  | 6077.626636 | 0.032076 |
| 210 | para | $\mathrm{A}_{1}$ | 2 | 0.000707 | 6288.146119 | 6288.1428 | -0.0033 | 6288.134723 | -0.011396 |
| 111 | ortho | $\mathrm{B}_{2}$ | 3 | 0.000637 | 6289.172875 | 6289.1739 | 0.0010 | 6289.128284 | -0.044591 |
| 121 | ortho | $\mathrm{B}_{2}$ | 4 | 0.000632 | 7420.092707 |  |  | 7420.077786 | -0.014921 |
| 102 | para | $\mathrm{A}_{1}$ | 1 | 0.002000 | 7576.381719 | 7576.3000 | -0.0817 | 7576.413281 | 0.031562 |
| 201 | ortho | $\mathrm{B}_{2}$ | 2 | 0.001414 | 7576.544710 | 7576.3000 | -0.2447 | 7576.596211 | 0.051501 |
| 300 | para | $\mathrm{A}_{1}$ | 2 | 0.001414 | 7752.263219 | 7751.9000 | -0.3632 | 7752.343205 | 0.079986 |
| 003 | ortho | $\mathrm{B}_{2}$ | 2 | 0.001414 | 7779.321260 | 7779.2000 | -0.1213 | 7779.352004 | 0.030744 |
| 131 | ortho | $\mathrm{B}_{2}$ | 1 | 0.002000 | 8539.561310 |  |  | 8539.565999 | 0.004689 |
| 112 | para | $\mathrm{A}_{1}$ | 2 | 0.001414 | 8697.141469 |  |  | 8697.133905 | -0.007564 |
| 211 | ortho | $\mathrm{B}_{2}$ | 2 | 0.001481 | 8697.154984 | 8697.3000 | 0.1450 | 8697.179341 | 0.024357 |
| 141 | ortho | $\mathrm{B}_{2}$ | 1 | 0.005000 | 9647.167310 |  |  | 9647.098855 | -0.068455 |
| 221 | ortho | $\mathrm{B}_{2}$ | 1 | 0.005000 | 9806.667310 |  |  | 9806.712978 | 0.045668 |
| 122 | para | $\mathrm{A}_{1}$ | 1 | 0.005000 | 9806.733119 |  |  | 9806.748170 | 0.015051 |
| 202 | para | $\mathrm{B}_{2}$ | 1 | 0.005000 | 9911.023119 | 9911.0500 | 0.0269 | 9911.102285 | 0.079166 |
| 301 | ortho | $\mathrm{A}_{1}$ | 1 | 0.005000 | 9911.023310 | 9911.0500 | 0.0267 | 9911.112478 | 0.089168 |
| 311 | ortho | $\mathrm{A}_{1}$ | 1 | 0.005000 | 11008.695310 | 11008.7800 | 0.0847 | 11008.774494 | 0.079184 |
| 302 | para | $\mathrm{A}_{1}$ | 1 | 0.001000 | 12149.460119 |  |  | 12149.552318 | 0.092199 |
| 104 | para | $\mathrm{A}_{1}$ | 1 | 0.015000 | 12524.637119 |  |  | 12524.834491 | 0.197372 |
| 401 | ortho | $\mathrm{B}_{2}$ | 1 | 0.015000 | 12525.214310 |  |  | 12525.346292 | 0.131982 |
| 312 | para | $\mathrm{B}_{2}$ | 1 | 0.015000 | 13222.762119 |  |  | 13222.790827 | 0.028708 |
| 213 | ortho | $\mathrm{A}_{1}$ | 1 | 0.015000 | 13222.772310 |  |  | 13222.798559 | 0.026249 |



Fig. 3. Deviations, in $\mathrm{cm}^{-1}$, between the MARVEL energy levels from this work and the variationally calculated linelist AYT2 [91]. Different colors represent different rovibrational symmetries (see Table 2 in Section 2.2).
formed as part of the ORBYTS project and further studies on other key molecules will be published in due course, involving highschool students from the UK, Australia, and Hungary. A paper discussing our experiences of performing original research in collaboration with school children can be found elsewhere [96].

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## Supplementary material

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jqsit.2018.07.012.

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