First detections of extragalactic SO$_2$, NS and NO

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Abstract. We report the first detections of SO$_2$, NS and NO in an extragalactic source, the nucleus of the starburst galaxy NGC 253. Five SO$_2$ transitions, three groups of hyperfine components of NO and five of NS were detected. All three species show large abundances averaged over the inner 200 pc of NGC 253. With a relative abundance of a few 10$^{-7}$, the emission of the NO molecule is similar or even larger than that found in Galactic star forming regions. The derived relative molecular abundances for each molecule have been compared with those of prototypical Galactic molecular clouds. These results seem to confirm that large scale shocks dominate the chemistry of these molecules in the nucleus of NGC 253, ruling out a chemistry dominated by PDRs for the bulk of the gas.

Key words. ISM: molecules – galaxies: individual: NGC 253 – galaxies: ISM – galaxies: starburst – galaxies: abundances

1. Introduction

Molecular lines from the central regions of galaxies are among the most promising tools to explore and understand the history of our universe (e.g. Combes et al. 1999 and references therein). In many galaxies most of the molecular emission stems from highly excited gas within the central few 100 pc. However, the processes driving the excitation and the complex chemistry of the gas are not entirely clear.

Molecular studies of nearby active galaxies such as NGC 253, M 82 or Arp 220 suggest that C shocks, photodissociating radiation, and irradiation by X-rays or cosmic rays (e.g. Rigopoulou et al. 2002; Güsten et al. 1981; Farquhar et al. 1994) play an important role in the heating and the chemistry of nuclear gas. The dominant mechanisms and their relative importance are still unclear.

Sulfur-bearing molecules such as SO$_2$ and NS are present in a wide variety of interstellar conditions, displaying enhanced abundances in the hot cores of high mass star forming regions. In hot cores, the chemistry of these molecules is determined by grain-mantle evaporation into warm gas (Charnley 1997). Abundance ratios such as NS/CS have been suggested to be one of the signatures of shocks in hot cores (Viti et al. 2001). The NO molecule is a fairly abundant molecule in the diffuse ISM of the Milky Way. This species is the main precursor of the N/O chemical network (Halfen et al. 2001).

In this letter we report the first detections of SO$_2$, NS and NO in an extragalactic source, the nucleus of NGC 253.

This region is one of the most prolific sources of molecular emission outside the Milky Way (Mauersberger & Henkel 1993). The large amount of hot gas (Mauersberger et al. 2003) and the high abundance of molecules with special chemistry, such as SiO (García-Burillo et al. 2000) indicates that the presence of large scale shocks should dominate the heating and the chemistry of the molecular gas. Also the very high abundance of the three molecules reported in this letter and their excitation can be explained if the chemistry of the nucleus of NGC 253 is dominated by large scale shocks.

2. Observations and results

Observations of the spectral lines (Table 1) were carried out with the IRAM 30 m telescope. Continuum measurements on nearby sources made every ~2 hours were used to keep a pointing accuracy better than ~3″. A wobbling secondary mirror was used, with a symmetrical beam throw of 4′ in azimuth and a switching frequency of 0.5 Hz.

As spectrometers we used two 256 × 4 MHz filterbanks for the 2 mm transitions and two 512 × 1 MHz filterbanks for the lines at 1.5 and 3 mm. The SIS receivers were tuned to SSB with an image band rejection >10 dB. Beam sizes were 21″ (at 3 mm), 17″ (at 2 mm) and 10″ or 12″ (at 1.5 mm). Spectra were calibrated with the standard dual load system. Intensities are given on a main-beam brightness temperature scale ($T_{MB}$). Linear baselines were removed from the spectra and the resulting profiles are shown in Fig. 1.
Fig. 1. Spectra of the SO$_2$, NS and NO transitions observed towards the nucleus of NGC 253 ($\alpha_{1950} = 00^h45^m06.0^s, \delta_{1950} = -25\degree33\arcmin45\arcsec$). Velocity resolutions are 10, 15 and 20 km s$^{-1}$ respectively. Ticks indicate the expected position of HF components of NS and NO. This parameter was fixed in the fit.

We detected five transitions of SO$_2$ in the 2 mm band, three groups of hyperfine components of NO and five of NS in the 2, 3 and 1.5 mm atmospheric windows. Profiles are consistent with the frequencies in the spectral line catalogues of Lovas (1992) and Pickett et al. (1998). We checked that no other known molecular transitions, both in the signal and the background, were consistent with the frequencies in the line catalogues of NS and NO. Table 1 shows the derived line parameters. The only line partially blended is SO$_2$ at 135 GHz that overlaps the nuclear emission of SO$_2$, NO and NS. According to measurements of CO and CN transitions with the 30 m telescope and SEST, we derive an extent of ∼200 pc at a distance of 2.5 Mpc; Mauersberger et al. 2003) This is in agreement with the size measured by Mauersberger et al. (2003) from the high angular resolution interferometric maps of CS $J = 2-1$ of Peng et al. (1996). Thus, we can obtain the source averaged brightness temperatures ($T_B = T_{MB} \frac{\theta_B^2}{\theta_{LSR}^2}$) over 20$''$ and derive source averaged column densities in the upper levels of the observed transitions assuming optically thin emission. In the case of NS and NO, this was done independently for each series. Figure 2 and Table 1 present the derived line parameters.

### Table 1. Parameters derived from Gaussian fit to SO$_2$ transitions and hyperfine structure fitting for NO and NS transitions.

<table>
<thead>
<tr>
<th>$\nu$ (MHz)</th>
<th>Transition $J - J'$</th>
<th>$T_{MB}$ (mK)</th>
<th>$V_{LSR}$ (km s$^{-1}$)</th>
<th>$\Delta v_{1/2}$ (km s$^{-1}$)</th>
<th>$T_M$ (mK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>134 004.8</td>
<td>$8(2,6) - 8(1,7)$</td>
<td>380 (90)</td>
<td>242</td>
<td>91</td>
<td>3.9</td>
</tr>
<tr>
<td>135 696.0</td>
<td>$5(1,5) - 4(0,4)$</td>
<td>1000 (200)</td>
<td>245$^a$</td>
<td>140$^a$</td>
<td>5.4</td>
</tr>
<tr>
<td>140 306.1</td>
<td>$6(2,4) - 6(1,5)$</td>
<td>650 (80)</td>
<td>248</td>
<td>115</td>
<td>5.3</td>
</tr>
<tr>
<td>146 605.5</td>
<td>$4(2,2) - 4(1,3)$</td>
<td>1080 (230)</td>
<td>241</td>
<td>148</td>
<td>6.9</td>
</tr>
<tr>
<td>151 378.6</td>
<td>$2(2,0) - 2(1,1)$</td>
<td>800 (200)</td>
<td>225</td>
<td>158</td>
<td>4.8</td>
</tr>
</tbody>
</table>

$^a$ Transitions and integrated intensity refer to the whole group of HF components while $T_{MB}$ and frequency refer to the main component.

3.2. Column densities and rotational temperatures

In order to derive reliable physical conditions when comparing line intensities measured with different beam sizes ($\theta_B$), we have to make assumptions on the overall extent ($\theta_{LSR}$) of the nuclear emission of SO$_2$, NO and NS. According to measurements of CO and CN transitions with the different beam sizes of the 30 m telescope and SEST, we derive an extent of the emitting region of ∼20$''$ (240 pc at a distance of 2.5 Mpc; Mauersberger et al. 2003) This is in agreement with the size obtained by Mauersberger et al. (2003) from the high angular resolution interferometric maps of CS $J = 2-1$ of Peng et al. (1996). Thus, we can obtain the source averaged brightness temperatures ($T_B = T_{MB} \frac{\theta_B^2}{\theta_{LSR}^2}$) over 20$''$ and derive source averaged column densities in the upper levels of the observed transitions assuming optically thin emission. In the case of NS and NO, this was done independently for each series. Figure 2

### 3. Analysis

3.1. Derived line parameters for SO$_2$, NS and NO

To derive the SO$_2$ line parameters, we have fitted a single Gaussian to the observed profiles. Figure 1 shows the Gaussian fits superposed on the observed spectra, and Table 1 presents the line parameters derived from the fit.

Fitting the NS and NO profiles is more complex as their ground state presents X-doubling. Thus each rotational level $J$ is split into two single rotational levels with opposite parity. We use the usual notation to label, in Table 1, the lower and upper series of single rotational levels as $e$ and $f$ for NS, and $\Pi^+$ and $\Pi^-$ for NO. Rotational levels are further split into hyperfine (HF) components (Gerin et al. 1992). Figure 1 indicates the location of the HF components, which are unresolved due to the broad intrinsic linewidth of the emission from the nucleus of NGC 253. Single Gaussian fits cannot be used in this case. We fitted the HF components simultaneously with a comb of Gaussian profiles with identical width and the relative frequency and line intensities fixed to the spectroscopic parameters of NS and NO. Table 1 shows the derived line parameters.
shows the population diagrams for each molecule where the level population can be described by a single rotational temperature represented as a solid straight line.

The derived $T_{rot}$ as well as total column densities and estimated fractional abundances are summarized in Table 2. To determine the abundance, we have assumed the H$_2$ column density of $1.7 \times 10^{22}$ cm$^{-2}$ derived by Mauersberger et al. (2003) from their $^{13}$CO measurements. The uncertainty of a factor of 2 in the H$_2$ column density (Mauersberger et al. 2003) does not critically affect the conclusions of this work. In general, the fractional abundances are high with the most outstanding case of NO with an abundance larger than $10^{-7}$. For the NS molecule, parameters derived for each series are: $N_{NS}(e) = 4.4 \times 10^{13}$ cm$^{-2}$, $T_{rot}(e) = 8$ K and $N_{NS}(f) = 6.4 \times 10^{13}$ cm$^{-2}$, $T_{rot}(f) = 7.6$ K. No substantial difference is found between series, in agreement with the result of Gerin et al. (1992) derived from NO in different Galactic molecular clouds.

### 4. Discussion

To establish the mechanism driving the chemistry in NGC 253, we compare in Table 3 the measured fractional abundances of SO$_2$, NS and NO with those in prototypical molecular Galactic clouds dominated by different types of chemistry. We have selected two dark clouds to illustrate the ion-molecule chemistry associated with quiescent gas, three hot cores to illustrate grain surface and shock chemistry associated to massive protostars and one photodissociation region (PDR) to illustrate the UV dominated chemistry produced by OB stars in the main sequence.

### Table 2. Source averaged column densities, rotational temperature and abundances.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>$N$ (cm$^{-2}$)</th>
<th>$T_{rot}$ (K)</th>
<th>$[X]/[H_2]^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SO$_2$</td>
<td>$7 \times 10^{14}$</td>
<td>14 (9)</td>
<td>$4 \times 10^{-7}$</td>
</tr>
<tr>
<td>NS</td>
<td>$5 \times 10^{13}$</td>
<td>8 (1)</td>
<td>$3 \times 10^{-9}$</td>
</tr>
<tr>
<td>NO</td>
<td>$5 \times 10^{15}$</td>
<td>7 (2)</td>
<td>$3 \times 10^{-7}$</td>
</tr>
</tbody>
</table>

$^a$ The uncertainty is a factor of 2 in both directions, due to the uncertainty of the assumed $N(H_2) = 1.7 \times 10^{22}$ cm$^{-2}$ (Mauersberger et al. 2003).

### Table 3. Fractional abundances compared with similar studies in molecular clouds.

<table>
<thead>
<tr>
<th>Source</th>
<th>$[SO_2]/[H_2]$</th>
<th>$[NS]/[H_2]$</th>
<th>$[NO]/[H_2]$</th>
</tr>
</thead>
<tbody>
<tr>
<td>NGC 253</td>
<td>0.4</td>
<td>3</td>
<td>30</td>
</tr>
<tr>
<td><strong>Dark Clouds</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L134N</td>
<td>0.4$^a$</td>
<td>0.2–0.6$^b$</td>
<td>20$^c$</td>
</tr>
<tr>
<td>TMC1</td>
<td>&lt;0.1$^e$</td>
<td>0.7–1.2$^e$</td>
<td>2.7$^e$</td>
</tr>
<tr>
<td><strong>Hot Cores</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orion Hot Core</td>
<td>9.4$^d$</td>
<td>0.4$^e$</td>
<td>30$^e$</td>
</tr>
<tr>
<td>Sgr B2(N)</td>
<td>3$^f$</td>
<td>10$^f$</td>
<td>20$^f$</td>
</tr>
<tr>
<td>Sgr B2(M)</td>
<td>40$^f$</td>
<td>0.03$^f$</td>
<td>30$^f$</td>
</tr>
<tr>
<td><strong>PDRs</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orion Bar</td>
<td>0.01$^g$</td>
<td>0.2$^e$</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Ohishi et al. (1992); $^b$ McGonagle et al. (1994); $^c$ Gerin et al. (1992, 1993); $^d$ Charnley (1997); $^e$ McGonagle et al. (1997); $^f$ source averaged Nummelin et al. (2001); $^g$ Jansen et al. (1995).

Chemistry dominated by PDRs can be ruled out for NGC 253 since the abundances of SO$_2$ and NO are much larger than those in Galactic PDRs. It is remarkable that only one source in Table 3, the hot core in Sgr B2(N), shows larger or similar abundances than those observed in NGC 253. This is surprising given the expected beam dilution in NGC 253 for molecules with a high dipole moment like NS and SO$_2$. However, it is very unlikely that hot cores dominate the emission of the three molecules since the low rotational temperatures derived are inconsistent with the high excitation temperature (>70 K) expected from the dense hot cores. This suggests that the origin of the large abundance of these molecules must be related to a hot core like chemistry, but in regions with moderate densities in which molecules are subthermally excited. Molecular clouds in the center of the Galaxy show hot core like chemistry produced by the ejection of molecules from the grains by large scale shocks (see e.g. Martín-Pintado et al. 2001). Chemistry driven by large scale C shocks has been proposed to explain the high temperatures and large abundances of NH$_3$ and SiO (García-Burillo et al. 2000; Mauersberger et al. 2003). We will now discuss in detail how the abundance of the molecules reported in this paper fits into this picture.
4.1. SO$_2$

Towards NGC 253, we find a SO$_2$ abundance between that found in dark clouds and hot cores. The relatively high rotational temperature of SO$_2$ rules out that the bulk of its emission arises from dark clouds with their typical kinetic temperature of 10 K. Within the context of hot core chemistry, SO$_2$ is rapidly formed from the H$_2$S evaporated from grain-mantles when $T < 200$ K. The same type of chemistry is expected if H$_2$S is injected into the gas phase by shocks. The observations of SO$_2$ towards Sgr B2 by Cummins et al. (1986) support this idea. They found that SO$_2$ emission arises from two components with very different rotational temperatures, the first one associated with the Sgr B2 hot core ($T_{\text{rot}} = 310$ K) and a second, subthermally excited one, with a much lower temperature of 26 K associated with the envelope. The excitation temperature is somewhat larger in the envelope of Sgr B2 than in NGC 253. However, the latter is averaged over a much larger region. We therefore conclude that SO$_2$ arises from the high temperature component observed in NH$_3$ by Mauersberger et al. (2003).

4.2. NO

The large NO fractional abundance averaged over an almost 200 pc region suggests that NO emission is fairly widespread through the whole nucleus of NGC 253. If this emission would arise from dark clouds, all the clouds in the nucleus of NGC 253 must be of the L134N type, contrasting with what we find in the solar vicinity where most of the clouds are like TMC1. However, the lack of SiO emission in dark clouds (Ziurys et al. 1989) suggests that large scale shocks are the origin of the NO abundances observed in NGC 253. Like in the center of our Galaxy, NO emission from NGC 253 could arise from warm moderately dense gas, such as that of the envelope of Sgr B2 which shows a particular chemistry due to strong shocks and/or irradiation by hard X-rays (Martín-Pintado 1997, 2000). It is worth noting that models of X-ray chemistry (Lepp & Dalgarno 1996) predict an enhancement of the NO abundance ($10^{-6} - 10^{-7}$) as observed in Sgr B2 and NGC 253. Both NGC 253 and the envelope of Sgr B2 have enhanced SiO and NO abundances suggesting a similar origin.

4.3. NS

The NS abundance in NGC 253 is higher than in most Galactic dark cloud cores and hot cores. McGonagle & Irvine (1997) studied the excitation of NS in several molecular clouds and found low excitation temperatures and moderate abundances in the hot and very dense regions as expected from the large destruction rate of this radical under hot core conditions ($n(H_2) > 10^6$ cm$^{-3}$, $T_k > 100$ K). Chemical models of hot cores indicate an enhancement of the NS relative abundance to CS in the presence of C shocks (Viti et al. 2001). The high NS/CS ratio of 0.4 (with X(CS) = 4 x 10$^{-9}$; Mauersberger et al. 2003) agrees with the prediction of these models in which a shock passed 10–20 x 10$^3$ years after the onset of radiative heating in the hot core. This favors large scale shocks as the most likely explanation for the NS abundance in NGC 253. It is interesting to note that this NS/CS ratio is an order of magnitude above the ratio measured by Hatchell et al. (2002) towards six hot cores.

The origin of the large scale shocks is not yet clear. High angular resolution images of nuclear SiO emission in NGC 253 suggest different origins (García-Burillo et al. 2000). In the inner circumnuclear disk, shocks from a dense molecular outflow driven by massive protostar dominates. In view of the low excitation temperatures derived for the new molecules, the bulk of the emission is probably not related to this origin. The other proposed causes, shocks in the outer Lindblad resonance as a consequence of a barred potential or generated by mass ejection from the disk, could explain the abundances and excitation temperatures.

Interferometric observations of SO$_2$, NO and NS are needed to measure their spatial distribution and to establish the origin of the large abundances of these molecules in the nuclear region of NGC 253.

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