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AN APPARATUS FOR ABSORPTION MEASUREMENTS IN THE FAR INFRARED BY FOURIER TRANSFORM

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Résumé. — L'article donne la description d'un système complet destiné à l'étude dans l'infra-rouge lointain de matériaux magnétiques à des températures aussi basses que 1,3 K et à des champs magnétiques allant jusqu'à 7,5 T. Les bonnes performances de l'ensemble sont illustrées par des résultats sur FeCl₂ et les ions de Fe²⁺ dans MgO.

Abstract. — The description of a complete system for far infrared studies of magnetic materials at temperatures as low as 1.3 K and magnetic fields up to 7.5 T is given. The good overall performance is demonstrated by the results on FeCl₂ and Fe²⁺ ions in MgO.

1. Introduction. — For absorption experiments in solids in the far infrared between 10 and 250 cm⁻¹ Fourier transform techniques offer certain advantages over a classical grating spectrometer:

— for a comparable measuring time a much better signal to noise ratio,
— direct wavelength calibration,
— precise knowledge of the instrumental profile.

In this wavelength range interferometers of the Michelson- or Twyman-Green type seem to be the most convenient instruments. One drawback with commercial interferometers is that, while working satisfactorily at low optical densities and samples at room temperature, they are normally not equipped for low temperature work and the standard detector they are supplied with, the Golay-cell, is not sensitive enough for experiments on dilute ions in a more or less absorbing matrix, e. g. Fe²⁺ in MgO.

So the necessary modifications and additions to such an instrument must take into account the following points:

— Sample temperature between 1.3 and 50 K.
— Low temperature bolometer such as Ge : Ga.
— Possibility to apply a magnetic field.

With these conditions in mind it becomes clear that the cryogenics part of the installation must be given particular thought.

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In this paper we wish to present the final stage of construction of an instrument conceived along such lines that has been operative now for several months.

As an example for the results obtained so far, we give the spectra of Fe²⁺ ions in MgO and the antiferromagnetic resonance (AFMR) mode of FeCl₂, a typical layered structure.

2. Interferometer. — We started out with a BECK- MAN/RIIC FS 720 module, which had already proved to be a very sturdy an mechanically sound instrument. The continuous mirror drive with a Moiré-frange system to monitor the path-difference was too wasteful of measuring time and not reliable enough. It was replaced by a stepping motor (SLO SYN SS25) driving directly the micrometer screw originally mounted in the instrument. A simple control unit was built generating repeatedly the appropriate measuring cycle:

— Rapid advance of the mobile mirror by a determined number of steps.
— Integrating period of several time-constants of the signal amplifier.
— Digitization and recording on punched tape.

In order to judge the quality of the lead screw and general positioning precision of the stepping motor we compared a water-vapour spectrum in the range between 40 and 150 cm⁻¹ with one published by Rao et al. [1] and found an agreement of the absorption frequencies better than 0.05 cm⁻¹, that is to say, better than the resolution of the spectra.

The source, a Philips HPK 125 high-pressure mer-

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cury arc, was originally choke-fed on AC. Its stability and longevity could be greatly improved by the use of a regulated current supply of 300 V, 2 A in series with a 100 Ω resistor. The long-term stability of the operating current (1 A) is of the order of $2 \times 10^{-4}$ A.

The light of the source passes a variable-frequency chopper permitting the use of a Golay-cell at typically 10 Hz as well as operation of a bolometer detector at its optimum frequency. This is achieved by feeding the coils of the modulator motor with two currents of variable frequency but a fixed phase difference of 90°.

3. Cryostat. — A cryostat has been built to meet the various requirements of far infrared spectroscopy of magnetic compounds such as FeCl$_2$ or isolated magnetic ions, pairs or triads in a non-magnetic matrix. It permits controlled sample temperature between 1 and 50 K and application of a magnetic field of up to 7.5 T in the vertical direction. It is further equipped with a Ge : Ga Bolometer working normally at 1.3 K controlled to better than 2.5 mK.

Sample and bolometer (Fig. 1) are both coupled to a pumped helium bath inside a long cylindrical enclosure, which is placed in liquid helium contained in a complete superinsulated Dewar. This cylinder is filled at ambient temperature with about $2 \times 10^3$ N m$^{-2}$ (15 torr) of helium. While cooling down the exchange gas pressure is maintained by heating the active-carbon cryo-pump. Upon arriving at 4.2 K the exchange gas is removed by turning off the heater current.

Liquid helium can be admitted into the inner thermostat via a needle-valve connecting it with the surrounding bath. This thermostat is pumped by a 20 m$^3$/h two-stage rotary pump and attains in the limit a temperature of about 1.03 K. The thermal loss of the Dewar is only 70 mW, thus giving an autonomy of 8 hours.

The superconducting solenoid may be fastened on the exterior of the sample enclosure with the bolometer well out of the magnetic field. With the magnet in operation the time between two top-ups necessary to keep the sorption pump covered by helium drops to about 3 h in the worst case of fields higher than 4 T and external feeding of the coil.

Light from the interferometer is picked up by a brass light-pipe at the focusing point in the former sample chamber. After passing a sealed TPX window, fixed in a Leybold-type quick-release coupling, it is guided down to the sample holder (Fig. 2) and the bolometer. The light-pipes in the cryostat are of thin-walled stainless steel to suppress thermal coupling between the outer helium, the sample holder whose temperature is variable between 1.1 and 50 K and the bolometer whose temperature is fixed at 1.3 K.

![Fig. 1. — Schematic view of the cryostat and bolometer housing.](image1)

The sample holder is shown coupled to the 4.2 K point, a copper rod fixed to the upper flange and passing through an insulated opening in the pumped helium can. It may alternatively be coupled to the latter by soldering the copper braid to a lug on its underside.

![Fig. 2. — Detail of the sample holders. The magnetic field is vertically oriented.](image2)

We use two types of sample holders (Fig. 2): One with a straight-through passage of the radiation for isotropic samples and layered structures with their c-axis parallel to the field direction. The other allows...
the c-axis of the layered samples to be oriented perpen-
dicularly to the field direction while keeping the wave
vector of the radiation parallel to the c-axis. This is
done by two mirrors at 45°. The resulting kink in the
light path is readily taken care of by inclining the light-
pipes. So the positions of sample holder and bolo-
meter housing do not change.

4. Bolometer. — As infrared detector we opted for
a Ge : Ga bolometer for its outstanding performances
in detectivity and N. E. P. The theory of this type of
bolometer has already been extensively treated by
several authors [2], [3] so we limit ourselves to an
outline of the preparation and the performance of the
bolometer we have mounted.

A slice of Ga-doped Ge was cut from a boule of
approx. 0.1 cm resistivity. After cutting and lapping
the sample was etched in mild alcohol etchant
(10 ml HNO₃ 60 % + 10 ml HF 38 % + 1 ml Etha-
nol + 18 ml dist. water). The final dimensions of the
bolometer are 6 × 10 × 0.1 mm³.

Ohmic contacts were obtained by copperplating two
corners of the chip in a classical acid copper-sulfate
bath and soldering with indium the copper leads of
0.1 mm diameter serving as electrical connection,
thermal impedance and mechanical support for the
bolometer. The leads, of a final length of about 10 mm,
have a thermal impedance of 55 µWdeg⁻¹. The bolo-
meter housing (Fig. 1), is coupled to the pumped He
bath by its suspending two copper tubes and is equipped
with a carbon thermometer and a heater resistance.

The value of the ambient resistivity of ~ 0.1 Ω cm
seems to indicate that the germanium used might be too
highly doped. However, preliminary measurements of
the R/T relationship yielded a value of the activation
temperature

\[ \frac{E}{k} = \frac{d(\log R)}{dT} \]

of \( \frac{E}{k} = 5.6 \) K at the temperature of 1.3 K, close to the
values reported by other authors [2], [3].

The responsitivity at the operating point of 1.3 K
was determined by the static method as used by pre-
vious authors [2], [4] which is based on on the V/I
relationship :

\[ S = \frac{(Z - R)}{2V}, \quad \frac{R_1}{R_1 + Z} = 40 \text{ kVW}^{-1} \]

where

\[ V \] is the voltage across the bolometer,
\[ R = V/I \] its resistance (427 kΩ),
\[ Z = dV/dI \] its dynamic resistance (362 kΩ) and
\[ R_1 \] the load resistance (10 MΩ metal-film at 4.2 K).

The thermal conductance is given by

\[ G = dP/dT = (dP/dR) \cdot (dR/dT) \]

and can be obtained through the P(I) plot and the
thermal measurement of R(T).

Since we are principally interested in the bolometer
as an IR-detector that is part of a complete exper-
imental setup for measuring absorption, we prefer to
evaluate its noise performance in conjunction with the
associated equipment such as biasing source and lock-
in amplifier.

The noise voltage thus understood, estimated by the
noise at the output of a PAR 128 lock-in amplifier was
better than 0.2 μV in a 1 Hz bandwidth. This gives
with the previous value for \( S \) a

\[ \text{NEP} = 5 \times 10^{-12} \text{ WHz}^{-1/2}. \]

The detector time-constant \( \tau \) was obtained by obser-
ving the variation of the phase angle with rising modu-
lation frequency. 45° is reached at about 40 Hz which
yields

\[ \tau = \frac{1}{2 \pi f(45°)} = 4 \times 10^{-3} \text{ s}. \]

We therefore adopted an operating frequency of

\[ f = 35 \text{ Hz}. \]

The following table lists again the pertinent charac-
teristics of the detector :

<table>
<thead>
<tr>
<th>Dim</th>
<th>6 × 10 × 0.1 mm³</th>
<th>( T )</th>
<th>1.3 K</th>
</tr>
</thead>
<tbody>
<tr>
<td>( I )</td>
<td>1.9 μA</td>
<td>( R )</td>
<td>427 kΩ</td>
</tr>
<tr>
<td>( R_1 )</td>
<td>10 MΩ</td>
<td>( d(\ln R)/d(1/T) )</td>
<td>5.6 K</td>
</tr>
<tr>
<td>( G )</td>
<td>55 × 10⁻⁶ WK⁻¹</td>
<td>( \tau )</td>
<td>4 × 10⁻³ s</td>
</tr>
<tr>
<td>( f )</td>
<td>35 Hz</td>
<td>( S )</td>
<td>40 kW⁻¹</td>
</tr>
<tr>
<td>( \text{NEP} )</td>
<td>5 × 10⁻¹² WHz⁻¹/₂</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

5. Conclusion. — The best way to show satisfac-
tory functioning of the apparatus as a whole is to give
samples of spectra actually measured with it.

![Graph](image)

**FIG. 3.** — Far infrared absorption spectrum of Fe²⁺ ions in MgO (0.43 % Fe) at 4.2 K. Resolution is about 4 cm⁻¹.

Figure 3 shows the absorption spectrum of Fe²⁺-ions in MgO (0.43 % Fe), an example of a dilute sys-
tem. The sample temperature was 4.2 K. The resolu-
tion is ~ 4 cm⁻¹, whereas the frequency range is
relatively large. The peak at 107 cm⁻¹ is due to the
(here unresolved) \( \Gamma_{3g} \rightarrow \Gamma_{3g} \) and \( \Gamma_{3g} \rightarrow \Gamma_{4g} \) transi-
tions of the splitted ground-state of the Fe-ion [5],
[6]. The absorption at 33 cm⁻¹ may be due to Fe-pair
interaction. The interferogram was taken with a 50 G
(0.012 mm) Mylar beamsplitter and sampled on
250 points up to the maximal path-difference of
± 0.25 cm. The recording took about one hour. Under
these conditions the smallest detectable value of the absorption coefficient for a sample of 1 cm thickness would be of the order of 0.1 cm$^{-1}$.

Figure 4 shows the AFMR-mode absorption in FeCl$_2$ which is situated on the lower side of our operating range (5-250 cm$^{-1}$). The interferogram of 1550 sampling points and a maximal optical path-difference of ± 6 cm was taken with a 400 G (0.1 mm) Mylar beamsplitter. The recording took about two hours using a 5 s integrating period. Since the thickness of the sample was 0.1 cm, the peak presents an absorption coefficient of 40 cm$^{-1}$. The position of the peak, 16.51 cm$^{-1}$, was reproducible to ± 0.03 cm$^{-1}$ on different runs. The resolution here is about 0.25 cm$^{-1}$ and we deduce that the transition has a natural width of 1.2 cm$^{-1}$.

Acknowledgments. — At this point we wish to thank O. Testard who designed the cryostat and J. Le Ny for assembling and testing the cryogenic equipment. We are further very grateful to A. Boix who constructed the mirror-control logic and the temperature regulators. The detector material has been contributed by B. Pajot with whom we had also many fruitful discussions on the subject.

References