Four hour identification of *Enterobacteriaceae* with the API Rapid 20E and Micro-ID systems

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**SUMMARY** One hundred strains of *Enterobacteriaceae* were examined in parallel with the API Rapid 20E and Micro-ID commercial four hour identification systems. With the API Rapid 20E system 78\% of the strains were correctly identified, 15\% were not identified, and 7\% were misidentified. The respective figures with the Micro-ID system were 74\%, 11\%, and 15\%.

The Micro-ID system and its predecessor Pathotec were among the earliest commercial systems designed to yield identification of *Enterobacteriaceae* after four hours of incubation. Later, API USA (Analytab Products, Plainview, New York), developed a data base to permit interpretation of the results obtained after incubating the standard API 20E system for only five hours (this data base is available only in America). More recently, API France (API System SA, La Balme-les-Grottes, Montalieu Vercieu, France, a company independent of API USA and which supplies the United Kingdom with API products) introduced a product specifically designed for four hour identification of *Enterobacteriaceae*, API Rapid 20E (this product was available in America as the DMS Rapid E System from DMS Laboratories, Flemington, New Jersey, but since October 1984 the product has been marketed by API USA as the API Rapid E system).

Only a few authors have performed an evaluation solely of the Micro-ID system. More often the Micro-ID system has been evaluated in parallel with another system, generally API 20E as an 18–24 hour identification system. In one of these studies the authors included time and motion and cost comparisons in their evaluation as well as success in identification. In addition, in two of these studies other commercial identification systems were included in the evaluation (AutoMicrobic System, Enterotube, and Micro-Media System, API 10S, Enterotube, Enterotube II, and Minitek). Some American authors have compared the Micro-ID system with API 20E, using the latter both as an 18–24 hour and as a five hour identification system. In two of these studies other commercial identification systems were also included in the evaluation (Abbott MS-2, Entero-Set 20). Gooch and Hill compared Micro-ID with API 20E, using the latter solely as a five hour identification system. Micro-ID has also been evaluated in parallel with the Minitek system and with the r/b system. Some workers have used the Micro-ID system for the direct identification of bacteria from blood (in two of these studies other commercial systems were included in the evaluation: Abbott MS-2, API 20E, and AutoMicrobic System, Minitek) whereas others have examined the effect of refrigeration of the inoculated system to permit delayed reading of the reactions. Cox et al modified the procedure for inoculating the Micro-ID kit, using a single colony to minimise the possibility of inoculating the kit with a mixed culture. The Micro-ID system has also been used for the rapid biochemical characterisation of *Haemophilus* species.

The API Rapid 20E system has been evaluated alone but more commonly in parallel with the standard API 20E system. In two of the latter studies other commercial identification systems were also included in the evaluation (Enterotube II and Mast ID-System, Abbott Quantum II). There have been two previous reports comparing the API Rapid 20E and Micro-ID systems, although in one of these studies the systems were used to directly identify bacteria from blood.

We report here our experiences obtained from an examination of 100 strains of *Enterobacteriaceae* examined in parallel with the API Rapid 20E and Micro-ID systems.

**Material and methods**

**ORGANISMS**

As only 100 units of each commercial system had been made available to us by the respective manu-
Four hour identification of Enterobacteriaceae with the API Rapid 20E and Micro-ID systems

facturers, only 100 strains could be used to assess the success of the API Rapid 20E and Micro-ID systems in identification (Table 1). The strains comprised a mixture of reference strains (maintained in the National Collection of Type Cultures) and field strains referred to the National Collection for identification. All the strains had been previously examined for 50 conventional characteristics and their identity confirmed or determined by the computer identification method of Lapage et al.42 and probability matrix of Bascomb et al.44 Although some field strains are referred to us because they are atypical in certain characteristics, others received for identification are typical but have been submitted for other reasons (such as test errors). Only fairly typical field strains were used in this evaluation.

**Tests**

All 100 strains were subcultured several times in an attempt to increase their enzyme activity. They were initially recovered on nutrient agar in Petri dishes. They were then allowed to grow for up to several days. This procedure was then repeated a further two times except that on the final subculture only overnight incubation was permitted (before inoculation of the commercial systems). All were then examined with the API Rapid 20E system, which consists of a series of plastic cups (containing dehydrated media) moulded to a strip of plastic. The following 21 tests were performed in the cups: β-galactosidase production (ONPG test), lysine decarboxylase, ornithine decarboxylase, urease production, citrate utilisation, deamination of

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Total 100 78 (12) 15 7 74 (1) 11 15

Numbers in parentheses are strains identified by reference to the full data base, either held on a floppy disc for use in a microcomputer (API Rapid 20E), or via the manufacturer (Micro-ID).

Escherichia aderacarboxylata is not specifically included in the data base of either product. Escherichia aderacarboxylata strains have, however, been considered to be biochemically atypical strains of Erwinia herbicola (synonym Enterobacter agglomerans). Strains of E aderacarboxylata should therefore either fail to identify on the system or identify to E herbicola; the E herbicola data base of the API Rapid 20E system does not allow for strains with the characteristics of E aderacarboxylata, unlike that of the Micro-ID. Serratia plymutica is not included in the data base of the Micro-ID system.
phenylalanine, malonate utilisation, aesculin hydrolysis, fermentation of arabinose, xylose, adonitol, rhamnose, cellulose, melibiose, sucrose, trehalose, raffinose and glucose, production of indole, production of acetoin, and production of cytochrome oxidase.

In accordance with the manufacturer’s instructions, the Rapid 20E kits were each inoculated with one well isolated colony (or several colonies of identical morphology where necessary) taken from overnight cultures grown on nutrient agar. The bacterial growth was removed from the plate using the tip of a polythene pipette (supplied with the strip). The bacterial growth was then homogenised in 1.25 ml of sterile physiological saline at 0.85% (and containing no additives) to achieve a turbidity equivalent to 0.5 on the McFarland scale (equivalent to 1 or 2 medium sized colonies). The tests were then inoculated with the suspension; in the citrate test two drops were added so as to fill both tube and lower part of the cupule, and in the other tests just sufficient inoculum was added to fill the tube section only. The lysine decarboxylase, ornithine decarboxylase, and urease tests were then overlayed with mineral oil before the tests were incubated for four hours at 37°C.

The reactions obtained were recorded after reference to a written description, provided by the manufacturer, of the appearance of positive and negative reactions. (A colour chart was not available at the time of this evaluation but one has since been produced; however, owing to limitations in printing inks such charts often offer no advantage over detailed written descriptions.) Reagents had to be added to the cupules testing for acetoin and indole production. The oxidase reaction of the strains had been determined previously by conventional means, but it may also be determined in the strip by the addition of the recommended reagent into the aesculin or phenylalanine cupules (if negative).

All the strains were also examined in parallel with the Micro-ID system, which comprises a moulded styrene tray containing 15 reaction chambers and a hinged cover. The first five reaction chambers contain a substrate disc and a detection disc. The remaining 10 reaction chambers each contain a single combined substrate/detection disc. These filter paper discs contain all substrate and detection reagents required to perform the indicated biochemical test (except that for acetoin production). The surface of the tray is covered with transparent polypropylene tape to contain the organism suspension during incubation and also provides complete visibility. The hinged cover is opened to provide access to the inoculation ports. The inside surface of the cover contains a strip of filter paper to absorb any spillage resulting from mishandling. The following 15 tests were performed in the reaction chambers: acetoin production, nitrate reduction, deamination of phenylalanine, hydrogen sulphide production, indole production, ornithine decarboxylase, lysine decarboxylase, malonate utilisation, urease production, aesculin hydrolysis, β-galactosidase production (ONPG test), and fermentation of arabinose, adonitol, inositol, and sorbitol.

In accordance with the manufacturer’s instructions, the Micro-ID kits were each inoculated with several well isolated morphologically identical colonies taken from overnight cultures grown on nutrient agar. The bacterial growth was removed from the plate using a wire loop and was then homogenised in 3.5 ml of physiological saline to achieve a turbidity equivalent to 2.0 on the McFarland scale (this is the inoculum density recommended for stock cultures; for fresh clinical isolates a turbidity equivalent to 0.5 on the McFarland scale is recommended). With the Micro-ID tray flat on the bench and the cover open the tests were inoculated with the suspension, about 0.2 ml per test. With the cover closed the Micro-ID tray was then incubated in an upright position for four hours at 37°C, after ensuring that the suspension was in contact with all substrate discs.

After incubation the Micro-ID tray was again placed flat on the bench and two drops of 20% KOH added to the acetoin test only. The tray was then held upright again to allow the KOH to flow down into the acetoin production test solution. Finally, the tray was rotated clockwise through 90° so that the upper (detection) discs in the first five wells became wet. The reactions obtained were recorded after reference to a written description, provided by the manufacturer, of the appearance of positive and negative reactions (a colour chart is available, however, for those who prefer it).

**IDENTIFICATION OF STRAINS**

The reactions of the 21 API Rapid 20E tests are recorded as a seven digit profile number based on octal numbers. This seven digit number is looked up in an index entitled *API Rapid 20E Analytical Profile Index* (product reference No 2079), and ours was dated March 1982. If a profile number was not listed in the profile index then the full data base, held on a floppy disc for microcomputer use, was consulted (the same information can be obtained by a telephone service offered by the manufacturer). In whichever way the data base is consulted a comment is given as to the level of acceptability of the suggested identification.

The reactions of the 15 Micro-ID tests are recorded in a similar way to yield a five digit profile.
number, which is looked up in an index entitled Micro-ID Identification Manual, and ours was dated January 1981. If a profile number was not listed in the index then the full data base was consulted through the manufacturer. For entries in the index a comment is given, as with API Rapid 20E, on the level of acceptability of the suggested identification.

For some identifications in both the API Rapid 20E and Micro-ID systems additional conventional tests, as well as serological investigations, may be required to further or confirm the identification. The additional tests were not carried out in this study; if the system indicated the correct taxon as one of the possible identities this was counted as a correct identification. If serological investigation would have prevented a misidentification then that pattern of results was counted as not identified.

Results

As can be seen from Table 1 the number of strains not identified was almost identical in both systems. API Rapid 20E had a higher identification rate (78%) than Micro-ID (74%). Without reference to the full data base, however, the identification rate with API Rapid 20E was only 66% compared with the corresponding Micro-ID figure of 73%. The misidentification rate with Micro-ID (15%) was twice that with API Rapid 20E (7%). Both systems encountered difficulty in identifying the strains of Citrobacter freundii, Providencia stuartii, Salmonella subgenus II, and Serratia liquefaciens. API Rapid 20E proved much more effective in identifying strains of Serratia plymuthica and Yersinia enterocolitica, but this is not surprising for the former taxon as S plymuthica is not included in the data base of the Micro-ID system. Conversely, Micro-ID proved more effective in identifying Escherichia aderboxyylata. This is not surprising either, however, for reasons given in the footnote to Table 1. For the remaining taxa the identification performance of the two systems was about the same.

Table 2 Misidentification of strains by the API Rapid 20E and Micro-ID systems

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*Not included in the data base of the Micro-ID system.

Discussion

In the few published evaluations of API Rapid 20E the identification success rate has generally ranged from about 90% to 96%. The lowest rates reported have been around 85%, although in one of these studies the identification rate rose to 92.4% when use was made of the full data base held by the manufacturer. Misidentification rates reported have been low, around 1.5%. The identification rate we obtained for the API Rapid 20E system (78%, which includes referrals to the full data base), is considerably lower than that reported by other authors. Similarly, the misidentification rate we observed (7%) was higher than that previously reported.

In the 30 or so published evaluations of the Micro-ID system the identification rate has been generally 90% or more, even when the kit was directly inoculated with blood or urine, and has sometimes exceeded 99%. A lower rate has been reported (87%) in a study in which the authors were inoculating the kit directly from blood cultures, and also in another study (74-2%) in which the author was working with atypical strains that could not be identified by conventional means. In the latter study, however, the author also examined the strains with the API Rapid 20E system and obtained a good identification rate (92.4%) despite the use of atypical strains. Misidentification rates reported with the Micro-ID system have varied between 1.5% for typical strains and 16-7% for atypical strains. The identification rate we obtained with the Micro-ID system (74%, which includes referrals to the full data base), is considerably lower than that reported previously. Other authors, however, had noted a misidentification rate (16-7%) in excess of that which we observed (15%), although using atypical strains.
identification rate, for both systems, is lower and our
misidentification rate higher than that reported by
most other authors. Firstly, we used only stock cul-
tures, and other authors\textsuperscript{5,8} noted lower identification
rates for such strains (for example, 93-2\% with
Micro-ID\textsuperscript{2}) than with fresh clinical isolates (for
example, 97-8\% with Micro-ID\textsuperscript{2}). This is because
stock cultures may show less enzyme activity than
fresh clinical isolates. Indeed, the manufacturer of
the Micro-ID system recommends a more dense
inoculum (approximately equal to a McFarland No 2
turbidity standard) for use with stock cultures. We
subcultured our strains several times in an effort to
increase their enzyme activity and increased the
inoculum density as recommended, but it is not possi-
able to judge how successful these procedures were.

The only way to test properly the effectiveness of
repeated subculturing would be to examine each
strain twice with the commercial system, once with
minimum subculturing and once with repeated sub-
culturing. The second possible reason for our lower
identification rate is that we chose a wider range of
organisms, including several rarer taxa, than would
normally be available to a routine clinical laboratory
performing such an evaluation. For example,
\textit{Escherichia coli} represented only 3\% of the strains
in our study, but in other studies this organism
accounted for 38-3\% (143/373)\textsuperscript{4} and 59\%\textsuperscript{14} of the
strains examined.

Whatever the factors that caused our
identification rate to be lower than that in most
other published evaluations of these products, those
factors apply equally to the two products. Therefore,
a comparison of the two, of which there are at pres-
et only two such reports,\textsuperscript{41,42} is of much greater
interest. From the procedures used it can be seen
that the API Rapid 20E product had several appar-
ent advantages over Micro-ID. Firstly, there were
21 tests as opposed to 15. With more tests, particu-
larly if wisely chosen, there can be greater discrimi-
nation between taxa so that in theory, at least, there
should be fewer instances of identification solely to
genus level and fewer instances in which supplemen-
tary tests need to be performed in order to further
the identification. Secondly, although both products
used the same density of inoculum (0-5 on the
McFarland scale) the volume of inoculum was con-
siderably greater (3-5 ml) with Micro-ID than it was
with API Rapid 20E (1-25 ml). To achieve sufficient
inoculum the Micro-ID kit will always require sev-
eral colonies, but with API Rapid 20E a single
colony will often suffice. This point will vary in impor-
tance with different workers, but most workers will
accept that with certain specimens, in particular
faeces, there is a greater danger that morphologi-
cally identical colonies may represent different
organisms. Others who have have inoculated the Micro-
ID system from primary isolation plates have found
that a sufficient inoculum density could, in any case,
be obtained only with a proportion of the cultures
(74\%,\textsuperscript{5,8} 87\%,\textsuperscript{8} 92\%,\textsuperscript{14} 83\%\textsuperscript{15}).

Since it may not always be convenient to read
Micro-ID tests after four hours several authors have
found that identification at the species level is essen-
tially unchanged when inoculated strips are refriger-
ated overnight and then incubated, or incubated and
then refrigerated overnight, before reading.\textsuperscript{30,32} It is
not yet known if it will be possible to do this with
API Rapid 20E. Finally, since there are fewer tests
with the Micro-ID system than with API Rapid 20E,
fewer combinations of test results are possible with
Micro-ID so the data base for that system is smaller
than that for API Rapid 20E. The Micro-ID profile
index is therefore more comprehensive than that of
API Rapid 20E. Consequently, fewer referrals need
to be made to the full Micro-ID data base (two
strains in this study) than to the full API Rapid 20E
data base (25 strains in this study). Such referrals
are, however, certainly worthwhile as the use of the
full data base increased the identification rate for
API Rapid 20E by 12\% in this study and 7-6\% in
another study\textsuperscript{14} (although the identification rate for
Micro-ID was increased by only 1\% in this study).

The identification rate of API Rapid 20E at 78\%
compared with that of Micro-ID at 74\% suggests
the former to be the more successful product, as
does the overall misidentification rate for API Rapid
20E (7\%), which is only half that for Micro-ID (15\%). The latter figures agree reasonably well with those of Altwegg\textsuperscript{41} (1-5\% and 16-7\%, respectively).

The proportion of strains not identified was about the
same with both systems.

The misidentifications with the API Rapid 20E
system (Table 2) should cause few serious conse-
quences. The \textit{Salmonella pullorum} strain that was mis-
identified as a \textit{Hafnia alvei} might cause problems.
Given the biochemical similarity of these two organ-
isms this problem could perhaps be avoided if, when
\textit{H alvei} was the suggested identification, the manu-
facturer recommended testing with a polyvalent
\textit{Salmonella} antiserum to exclude the possibility of a
misidentified \textit{Salmonella}. Other authors have noted
as we have, that occasional strains of \textit{Citrobacter freundii} may be misidentified as \textit{Escherichia coli}\textsuperscript{39}
and occasional strains of \textit{Serratia liquefaciens} may be
misidentified as \textit{Serratia odorifera}\textsuperscript{37} with the API
Rapid 20E system.

The misidentifications with the Micro-ID system
(Table 2) were confined almost exclusively to the
genera \textit{Klebsiella}, \textit{Providencia}, \textit{Serratia}, and \textit{Yer-
sinia}. With \textit{Providencia} and \textit{Yersinia} only was the
species identification incorrect, and a similar obser-
Four hour identification of Enterobacteriaceae with the API Rapid 20E and Micro-ID systems

Four hour identification of Enterobacteriaceae with the API Rapid 20E and Micro-ID systems


Cox NA, Bailey JS, Thomson JE. Comparison of Micro-ID and Minitek-serology systems for rapid identification of Sal-

References


2. Cox NA, Mercuri AJ. Accuracy of Micro-ID for identification of Yersinia enterocolitica,7 Klebsiella pneumoniae as Serratia marcescens as Hafnia alvei.23 and Yersinia enterocolitica as Yersinia pseudotuberculosis.2 Stuver and Matsen identified only nine of 16 strains of Providencia stuartii using the Micro-ID system, and others have considered the product inaccurate for identifying Providencia as well as Serratia.16 Others have noted the misidentification of Klebsiella strains to Serratia21 and vice versa, and this is in accord with our own findings.

Some misidentifications are inevitable with commercial identification systems because a lower threshold identification level is used than in the reference laboratory situation. The overall misidentification figures suggest that the data base for Micro-ID is more lenient than that for API Rapid 20E. Potentially weak areas in the data bases of both products might warrant the attention of the respective manufacturers.

Prices can change relatively quickly and the accuracy and ease of use of a commercial identification system are likely to be given greater weight than cost, but for those interested, the current price for API Rapid 20E is £1.38 compared with £1.28 for API 20E. Since the two prices are so similar, a major factor influencing which system to use will be whether one wishes to identify a strain in four hours or 18 to twenty four hours. The unit cost for Micro-ID is £2-70, which is almost twice that of API Rapid 20E. Both companies offer discounts on bulk purchases, which will reduce the above prices by up to 10-15%.

We thank API Laboratory Products Limited and General Diagnostics for their respective gifts of API Rapid 20E and Micro-ID kits.


Four hour identification of Enterobacteriaceae with the API Rapid 20E and Micro-ID systems.
B Holmes and P S Humphry

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