Long-term Programme for Pollution Monitoring and Research in the Mediterranean Sea (MED POL Phase II)

METHODS FOR MONITORING SELECTED POLLUTANTS IN SEWAGE EFFLUENTS AND COASTAL RECREATIONAL WATERS

Report on a joint WHO/UNEP Meeting

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PREFACE

During the pilot phase of the Mediterranean Pollution Monitoring and Research Programme (MED POL Phase I), carried out between 1976 and 1981, the project on Coastal Water Quality Control (MED POL VII), jointly coordinated by WHO and UNEP, was concerned primarily with bacteriological and related parameters for the monitoring of coastal recreational waters, shellfish-growing waters and shellfish flesh. During the course of this pilot project, in which thirty Mediterranean laboratories participated, the two principal methods of bacteriological analysis utilized were the membrane filtration culture (MF) technique and the most probable number (MPN) technique. A comprehensive intercomparison of these "wo methods was not finalized during the period of the pilot project.

Concurrently, during this period, work was started by the United Nations Environment Programme (UNEP) in cooperation with the World Health Organization on the preparation of standard reference methods for bacteriological and related sampling and analysis. These methods are designed for use by Mediterranean laboratories participating in the Long-term Programme for Pollution Monitoring and Research in the Mediterranean Sea (MED POL Phase II). This programme is being undertaken by Mediterranean coastal states as part of the Mediterranean Action Plan under the terms of the Convention for the Protection of the Mediterranean Sea against Pollution and its related protocols. These methods form part of a comprehensive series being prepared by UNEP's Regional Seas Programme Activity Centre (RS/PAC) in collaboration with the relevant United Nations Specialized Agencies, designed not only to cover all the possible parameters listed in the annexes to the Convention and protocols, but also to enable their utilization in regions other than the Mediterranean.

During the first half of 1982, four reference methods, covering the major microbiological indicator organisms for monitoring of coastal recreational waters, shellfish-growing waters and shellfish flesh, reached their final draft stage, prior to testing, review and revision. These methods were as follows:

- determination of total coliforms in seawater by the membrane filtration culture method;
- determination of faecal coliforms in seawater by the membrane filtration culture method;
- determination of faecal streptococci in seawater by the membrane filtration culture method;
- determination of faecal coliforms in bivalves by the most probable number method.

A number of Mediterranean laboratories were requested to test these methods under their local environmental conditions. Following this, representatives of such laboratories were invited to participate in a joint intercalibration exercise on these methods, which was held at the Istituto Superiore di Sanità, Rome, on 22-23 November 1982.

In order to facilitate the eventual application of the reference methods to other regions, representatives from three institutions outside the Mediterranean (two from the Kuwait Action Plan region, and one from the Caribbean region) were also invited to participate in the joint intercalibration exercise.

The first draft of a further reference method "Guidelines for monitoring the quality of coastal recreational and shellfish-growing waters" was also completed during the second half of 1982.

Early in 1982, a number of Mediterranean laboratories (other than those participating in the testing and intercalibration of the four microbiological reference methods) were invited to participate in an exercise on comparative testing of the membrane filtration culture (MF) and most probable number (MPN) methods, as well as to carry out determinations of supporting parameters. This particular study was carried out in cooperation with the Istituto Superiore di Sanità, Rome, which also undertook the preparation of the necessary synthesis and evaluation of results obtained. This study covered pollution sources (sewage effluents) as well as coastal recreational waters and reference areas.

The main objectives of the present consultation meeting were the following:

- to review the results of the comparative study on the MF and MPN methods undertaken by the different laboratories;
- to review the results of the study on sampling and analytical techniques utilized on pollution sources (sewage effluents), coastal recreational waters and reference areas;

- to review the results of the testing of microbiological reference methods carried out by Mediterranean laboratories, as well as those of the joint intercalibration exercise, to enable identification of technical and other problems, including quality control, and in the light of these, to review the relevant reference methods with a view to their confirmation or proposed revision:
- to make appropriate recommendations on sampling and analytical techniques on the parameters in question, vis-à-vis the Long-term Mediterranean Pollution Monitoring and Research Programme (MED POL Phase II) and other such regional activities.

The meeting was organized by WHO and UNEP in collaboration with the Istituto Superiore di Sanità, Rome.

Representatives of institutions participating in (a) the study on sampling and analytical techniques, including the intercomparison of the MF and MPN methods, and (b) the testing and joint intercalibration of bacteriological reference methods, as well as a number of selected international experts in the field both within and outside the Mediterranean region, were invited to attend the consultation meeting. In addition, the following international organizations and agencies were invited to send representatives: Food and Agriculture Organization (FAO), Intergovernmental Oceanographic Commission (IOC), United Nations Educational, Scientific and Cultural Organization (UNESCO), World Meteorological Organization (WMO) and International Atomic Energy Agency (IAEA).

Opening of the meeting (agenda item 1)

The meeting took place at the Istituto Superiore di Sanità, Rome, from 24 to 26 November 1982. It was attended by 28 temporary advisers from the Mediterranean and other regions. Ten Mediterranean and five non-Mediterranean countries were represented. There was one representative from UNEP and one from WHO/EURO. A list of participants is given at Annex 1.

Dr L.J. Saliba, Senior Scientist for the Mediterranean Action Plan, Regional Office for Europe of the World Health Organization, opened the meeting on behalf of the Regional Director, Dr Leo A. Kaprio. He referred to the current meeting, and to the two exercises within the framework of which it was convened, as the natural continuation and follow-up to the pilot project on coastal water quality control (MED POL VII) implemented between 1976 and 1981. He also expressed WHO's appreciation of the work undertaken and the facilities provided by the Istituto Superiore di Sanità, both in connexion with the consultation meeting itself and the joint intercalibration exercise, as well as in the study on sampling and analytical methods. He mentioned the particular appropriateness of this meeting being convened at the same venue as the series of meetings held throughout the course of the MED POL VII pilot project.

Professor L. Villa, Director of the Department of Environmental Microbiology, Istituto Superiore di Sanità, welcomed the participants on behalf of Professor F. Pocchiari, Director of the Institute. He expressed the Institute's appreciation at being able to continue its long-standing collaboration with WHO in this important field, and augured a fruitful and successful meeting. Referring to the exercise on sampling and analytical methods, he stated that the response by participating institutions was very encouraging. A wide range of topics had been covered, and valuable data obtained, although the range of contributions and methodologies employed had rendered the process of synthesis and comparability rather difficult.

Scope and purpose of the meeting (agenda item 2)

Dr L.J. Saliba explained the scope and purpose of the meeting. He described the general situation existing at the termination of the pilot project on coastal water quality control (MED POL VII) in 1981. Among the outstanding matters, he referred to the need for a comprehensive intercomparison of the MF and MPN methods, as well as for the development of methods for the sampling and analysis of supporting parameters. In this connexion, the study undertaken by the Istituto Superiore di Sanità, in which various Mediterranean laboratories had collaborated, fulfilled the former need, and the results would be reviewed during the current meeting. Regarding the reference methods, it was not within the scope of the current meeting to adopt a final revision. The methods would be reviewed as comprehensively as possible within the permissible time limitations. Comments and suggestions by participants would be recorded and fully taken into account in the preparation of revised versions of the methods, which would then be submitted to the Governments of Mediterranean States for use in national monitoring programmes. Even these revised versions could not really be described as final, as they would be continually updated as and when necessary.

Mr F.S. Civili, Marine Scientist, UNEP Coordinating Unit for the Mediterranean Action Plan, spoke both on behalf of the Coordinating Unit and of the Regional Seas Programme Activity Centre. He stated that in the coordination of the scientific components of the various Regional Seas programmes, including the Mediterranean Action Plan, UNEP had always given priority to the harmonization of analytical methodologies used by participating laboratories. In the MED POL programme, particularly during the pilot phase, a number of common sampling and analytical methods were recommended, and an intercalibration exercise covering some mandatory parameters organized. This was done in order to assist laboratories in the acquisition of comparable data necessary for

MED POL was now in its second phase, and monitoring had become mandatory for all Mediterranean countries. This meant that data being collected should now be fully reliable and comparable in order to allow UNEP to assess the state of pollution of the Mediterranean. As a result, to ensure comparability of data on a worldwide basis, thus contributing to the Global Environment Monitoring System (GEMS) of UNEP, reference methods and guidelines for marine pollution studies were being developed and recommended to governments for adoption. These methods and guidelines were being prepared by UNEP in cooperation with the relevant specialized bodies of the UN system and were being tested by a number of experts competent in the particular fields. The methods being discussed during this meeting had been developed in cooperation with WHO.

Finally, it was the intention of UNEP to strengthen the intercalibration component of the MED POL Phase II programme as well as in equivalent programmes carried out in other regional seas, and through this, to achieve strict quality control of all data obtained. For this purpose, similar exercises would be organized in future both in the Mediterranean and in other regions.

Election of officers (agenda item 3)

Professor L. Villa was elected Chairman, Dr S. Sotiracopoulos Vice-Chairman, and Mr V. Gauci Rapporteur. Dr L.J. Saliba acted as Secretary to the meeting.

Adoption of the agenda (agenda item 4)

The provisional agenda was unanimously adopted.

Review of the exercise on sampling and analytical methods (agenda item 5)

Reports were presented by a number of participants, following which a summary of data received prior to 15 October 1982 was also given, as prepared and processed by the Istituto Superiore di Sanità (appearing as Annex 1 to this report). Reports by participating institutions are given at Annexes 2 to 8. From these reports and the ensuing discussion, the following points emerged.

5.1 Number of samples

Since any laboratory participating in a monitoring programme can of necessity only cope with a limited number of samples (actual maximum depending on individual resources available), it should be decided which of the following alternatives was the best: (a) to examine a small number of samples in duplicate and triplicate, or (b) to examine more samples without duplication. Since the state of pollution of a marine ecosystem cannot be assessed on the basis of only a few samples, it is often acceptable to sacrifice duplication for the sake of examining more samples.

5.2 Examination of sand and sediment

It was considered of the utmost importance that recommended methods be simple and explicit. Otherwise results may not be comparable. This was particularly stressed with regard to the examination of sand and sediment. Such factors as sampling procedures, nature of rinse water, time of rinse and analytical procedure used on rinse water must be standardized. Various techniques for transferring the organisms from the sand particles to the water phase were mentioned. These included the use of blending machines, ultrasonic waves and addition of enzymes to rinse water.

5.3 Effects of method on organisms' survival

The point was also raised that some of these methods could potentially damage the organisms. Light is a very important factor in the die-off of indicator organisms. It was reported that a close correlation exists between bacterial indicator numbers in natural water and such factors as duration of daylight and time of day. Such actors should, therefore, be recorded during sampling It is important not only to stan ardize analytical techniques but also the processing of results.

5.4 Stressed organisms

The presence of stressed organisms in the marine environment was considered. Resuscitation techniques, such as pre-incubation of membrane at $36\,^{\circ}\text{C}\pm1\,^{\circ}\text{C}$ for faecal coliforms, was reported to give higher counts than using $44.5\,^{\circ}\text{C}\pm2\,^{\circ}\text{C}$ incubation temperature from the start. It was also proposed to recommend a holding medium in the case of sampling points which are far from the laboratory.

Stress involves any alteration in the normal metabolic processes of the cell, including degradation of genetic material and cytoplasmic constituents. Stress factors towards indicator organisms, e.g. unfavourable temperature, fluctuating temperatures, toxic substances and competition with a large number of organisms, do occur in the natural environment. The result of stress could be a temporary loss of biochemical characteristics or a prolongation of the lag phase of growth. Stressed organisms could affect negatively both MPN and MF techniques: a faecal coliform may fail to produce a recognizable colony on the membrane, or gas with the MPN technique. Faecal streptococci are particularly subject to stress. This could be due to their longer persistence and therefore contact with stress factors. Results of comparative experiments using primary cultures with and without resuscitation showed that the adoption of a resuscitation procedure in the isolation of these indicator organisms from natural water was highly recommendable. A number of laboratory procedures accentuate stress, e.g. primary culture in a inhibitory medium and high temperature. These must be avoided as much as possible. It is important to improve the recovery of stressed indicator organisms since stressed pathogens may be as viable and virulent as non-stressed pathogens. A number of resuscitation techniques, some of which can be used in combination, were proposed for further study and subsequent adoption.

- (1) Primary incubation at 36°C±1°C for two hours followed by incubation at 44.5°C±0.2°C.
- (2) Primary culture on a non-selective medium, applicable to both MPN and MF techniques. In the former the presumptive test is preferably carried out in Lactose Broth with subsequent subculture onto Mac Conkey Brilliant Green Bile Broth. In the MF technique, the membrane may be initially cultured on a nutrient agar and later transferred to selective medium.
- (3) The layered plate technique is applicable to MF. The membrane is cultured on a layered agar medium made up of a lower layer of selective agar medium overlain with a thin layer of nutrient agar. During the first two hours of incubation there occurs a slow diffusion process of the two agar layers.

During the discussion of attenuated organisms and their recovery it was pointed out that resuscitation occurs at the expense of selectivity and that in practice a compromise must be sought.

5.5 Media

Agar media were to be preferred to absorbent pads soaked with liquid media for the MF technique. It was pointed out that certain batches of absorbent pads contain toxic material and that the agar media are more convenient to use since they can be prepared and poured as needed, stored, and used when required, while pads saturated with liquid media must be used the same day they are prepared.

5.6 Time-lag between sampling and analysis

It was agreed that in all cases this time-lag must be kept to a minimum, and preferably within a total of 8 hours. During transport to the laboratory, the samples should be kept chilled (10°C) and protected from light, since it is known that the die-away rate in the presence of light can be very high indeed. It was suggested that no more samples are taken than can be analysed during the same day. When this is impossible the samples should be refrigerated (4°C) and analysed not more than 24 hours later. One particular technique of processing sewage samples after 24 hours was reported. This consisted in diluting the sample 1 in 10 with phosphate buffer prior to refrigeration. It was reported that the coliform concentration in the original sample processed on the same day (i.e. when it was taken) correlated very well with the diluted refrigerated sample tested on the following day.

5.7 Comparison of MPN and MF techniques

The comparative investigation of the MPN and MF techniques carried out showed good correlation between data obtained by MF with regard to faecal coliforms at the critical level, i.e. between 100 and 2000 faecal coliforms per 100 ml. It was noted, however, that it would be necessary to study further the correlation at the concentration which is significant for deciding whether a particular water body is fit for recreational or shellfish-growing use. The choice of methods was

therefore to be based on other considerations, in particular legal aspects, nature of samples and cost. With regard to the first, it appeared that for most participating countries, existing legislation recognizes both methods of analysis. With regard to the second, it was agreed that for turbid samples, the MPN was the method of choice. The same held good for highly contaminated samples, since it was anomalous to carry out a high dilution and then to concentrate the organisms on the membrane. With regard to cost, the situation is more complicated. MPN is more labour-intensive than MF. Since labour costs differ from country to country, it was necessary to carry out a costing exercise on the two methods in each and every country before taking a final decision. In this respect reference was made to a costing exercise carried out by Professor Geldreich in the USA. This was a very detailed exercise and could serve as a model on which to base similar exercises in other countries. During the discussion, the point was mentioned that re-use of plastic Petri dishes and even membranes could be considered in emergency situations.

6. Review of the results of the joint intercalibration exercise on reference methods (agenda item 6)

The meeting reviewed the results of the joint intercalibration exercise on reference methods carried out in the laboratories of the Istituto Superiore di Sanità, Rome, on 22-23 November 1982. A description of this exercise, giving background, participating laboratories, methodologies and results, including statistical analysis, is given at Annex 9.

It was agreed by the meeting that this exercise had proved most useful, and it was considered that such exercises were essential within the framework of the monitoring programme.

7. Review of draft reference methods (agenda item 7)

The meeting discussed and reviewed the draft reference methods for determination of bacteriological parameters (documents ICP/RCE 211(2)/7, 8, 9, 10 and 11). Prior to the general discussion, participants reported on the results of tests carried out under local environmental conditions, using these methods. Written reports were also submitted by the Environmental Health Laboratory, Hadassah Medical School, Jerusalem, and the Laboratoire départemental de Santé publique, Marseille. As a result of the discussions, a number of amendments to the draft reference methods were proposed.

7.1 Reference method No. 1 - Guidelines for monitoring the quality of coastal recreational and shellfish-growing waters

The draft of this particular reference method was still at an early stage of formulation and new substantive material was still being introduced. It was suggested that participants could more appropriately limit their comments to general suggestions on improvements to the present text and content of the final version, although specific suggestions would also be welcome. It was understood that the relevant parts of the text would be modified to cover requirements for shellfish-growing waters, and that evaluation of results (i.e. statistical analysis) would also be catered for, either in the guidelines themselves, or separately. Important points of the discussion which followed were:

Definition of sampling sites

Sampling should be carried out in zone of contact. Thus for water, a depth of up to 1.5 m was considered best. Also suggested were sand and sediment analysis. It was pointed out that contact with sand (sun-bathing, etc.) was usually longer than contact with water. Sediments, and especially the sediment/water interphase, were of special importance. More viruses and Salmonella could usually be isolated than from the overlying water. Special methods have been developed in the USA for the examination of the air/water interphase.

Monitoring of organisms other than the indicator species

Various micro-organisms besides total coliforms, faecal coliforms and faecal streptococci are of importance and perhaps deserve more attention than has been given to them up to now. These include pathogenic organisms. It was suggested that fungi may be better parameters to monitor recreational waters in tropical and subtropical regions.

Parameters to be monitored

It was noted that in the long-term MED PrL monitoring programme, there is only one mandatory microbiological parameter: faecal coliforms. It was made clear, however, that this by no means suggests that other parameters are of no value and may not be investigated. It was pointed out that faecal coliforms are a good parameter by which to monitor enteropathogenic bacteria while

faecal streptococci may be considered as a corroborating parameter in this respect. Faecal streptococci are known to persist longer in the marine environment so that they may be considered as indicators of a more remote pollution. Moreover, Streptococcus faecalis var. liquefaciens is known to be ubiquitous in nature and is discharged in great numbers by food industries. A close correlation between faecal streptococci and enteropathogenic viruses was reported for seawater. Higher counts of faecal streptococci are obtained using KF-medium than by using M-enterococcus-medium, both incubated at 36°C. This was attributed to a better recovery of enterococci originating from farm animals. Since this is known to be an important source of human pathogens, it was considered essential that indicators which are known to be coming from this source be also monitored. In this respect KF-medium is, therefore, superior to M-enterococcus-medium.

7.2 Reference method No. 2 - Determination of total coliforms in seawater by the membrane filtration culture method

The following amendments were proposed:

(1) an air incubator at 36°C±1°C to be used;

(2) M-endo-agar medium to be used. After 24 hours' incubation, red to pink colonies with a golden metallic sheen are counted on membranes having 20 to 80 total colonies. Other colonies need not be counted;

(3) confirmation not systematic but only initially and when in doubt;

(4) Mac Conkey Broth and BGB Broth to be used in the confirmation. Incubation is at 36°C±1°C for 48 hours;

- (5) as regards expression of results, this should be simplified by just rounding off counts to two significant figures;
- (6) a stock culture may be used for the Precision Estimation Test.

7.3 Reference method No. 3 - Determination of faecal coliforms in seawater by the membrane filtration culture method

The following amendments were proposed:

M-FC-agar to be used;

- (2) rosolic acid to be used only when interference by non-coliform organisms is experienced;
 (3) 5 cm Petri dishes to be used. These should be incubated in rigid metal or plastic watertight containers totally immersed - or water bath such that temperature of incubation is 44.5°C±0.2°C;

after 24 hours' incubation, blue colonies are counted on membranes having 20 to 80 total colonies. Non-blue colonies need not be counted;

(5) confirmation need not be systematic but carried out only at the start to be sure that what one is counting is really faecal coliforms. Thereafter, confirmation to be carried out when in doubt regarding the identity of organisms as judged by the colony morphology;

(6) confirmation to be carried out by subculture onto Mac Conkey Broth or Brilliant Green

Bile Broth, incubation being at 44.5°C±0.2°C for 24 hours;

(7) as regards expression of results, this should be simplified by just rounding off counts to two significant figures;

(8) a stock culture may be used for the Precision Estimation Test.

7.4 Reference method No.4 - Determination of faecal streptococci in seawater by the membrane filtration culture method

The following amendments were proposed:

an air incubator at 36°C±1°C to be used;

(2) KF-agar to be used. The medium must be boiled first to dissolve agar, left to cool to about 50°C, sterilized by filtration through membranes of pore size 0.2 um, TTC added and poured. The TTC indicator is heat and light labile;

(3) as regards expression of results, this should be simplified by just rounding off counts

to two significant figures;

- (4) a stock culture may be used for the Precision Estimation Test.(5) confirm if necessary.

7.5 Reference method No.5 - Determination of faecal coliforms in bivalves by the MPN method

The following amendments were proposed:

- (1) Lactose Broth incubated at $36^{\circ}\text{C}\pm1^{\circ}\text{C}$ to be used for presumptive test. Positive tubes are subcultured onto Mac Conkey Broth or BGB Broth and tryptone water medium and incubated in water bath at $44.5^{\circ}\text{C}\pm0.2^{\circ}\text{C}$;
- (2) phosphate buffer or 0.1% peptone water to be used as diluent;(3) simplify calculation with regard to subdivision of sample;
- (4) rewrite MPN table, so that it refers to grams of sample and not to volumes.

It was agreed that the next draft of reference method No.1 should reflect the general and specific points mentioned in the discussion. It was similarly agreed that the final versions of reference methods 2, 3, 4 and 5 should incorporate the amendments proposed. An outline summary of these four methods as finally agreed on by the meeting appears at Annex 10.

8. Future action and recommendations (agenda item 8)

The meeting discussed health-related monitoring within the framework of the long-term phase of MED POL. Apart from the specific recommendations made in connexion with the various agenda items above, the following recommendations were also made by the consultation meeting:

- (1) In the research component of MED POL, activities to be carried out should include (a) development of methods in connexion with pathogenic organisms, and (b) studies on factors which could affect results, i.e. sampling times, etc., in the case of indicator organisms currently being monitored.
- (2) Quality assurance studies are essential and it is important to ensure that results obtained are comparable. For this purpose it is necessary to carry out periodic intercalibration exercises. Such exercises, on the lines of the one organized in Rome on 22-23 November 1982, should be held in centralized laboratories on both an intra-country and an intercountry basis. Participants should work completely on their own, but each count should be made by more than one participant.
- (3) Apart from such exercises, standard cultures of organisms and media could also be sent to laboratories, and participants requested to make three or four determinations.
- (4) Studies should be carried out on the relation between bacterial concentration and the spread of disease. In this connexion, it was recognized that such studies are difficult because of the many factors involved.
- (5) Precise beach surveillance and control has to be included together with microbiological monitoring.
- (6) More frequent meetings of investigators participating in the programme should be held. In this connexion, the meeting considered that one of the most significant factors contributing to the success of the MED POL VII Pilot Project was the convening of regular meetings of principal investigators.

Annex 1

A STATISTICAL ANALYSIS OF THE RESULTS OF THE INTER-LABORATORY EXERCISE ON SAMPLING AND ANALYTICAL METHODS

by L. Villa, F. Aulicino, A. Piccioni and G.A. Zapponi Istituto Superiore di Sanità, Rome, Italy

1. Introduction

This report is a statistical analysis of the data received from Mediterranean institutions participating in the inter-laboratory exercise on the comparison of the membrane filtration (MF) and the most probable number (MPN) methods in determination of bacterial concentrations in coastal water quality monitoring.

Reports were received from the following institutions:

- (1) Institut national de la Santé et de la Recherche médicale, Nice, France
- (2) The A. Felix Public Health Laboratory, Tel-Aviv, Israel
- (3) Institut Pasteur de Tunis, Tunisia
- (4) Rudjer Boskovic Institute, Rovinj, Yugoslavia
- (5) Istituto Superiore di Sanità, Rome, Italy.

The following data are summarized in Table 1:

- sampling periods
- sampling points
- total number of water, sand and/or sediment samples analysed by each institution
- parameters monitored
- parameters proposed prior to the commencement of the exercise.

Data concerning the methods utilized by each participating institution are summarized in Table 2. In this context, a number of difficulties arose in the comparison of data collected, due to differences in incubation temperature, incubation time and culture media utilized. Detailed analyses of the method and data interpretation techniques used by individual laboratories are given in the relevant reports.

Methodology

In the statistical treatment of material, only quantitative data were considered. In particular, sets of data corresponding to specific localities and data including one or more values expressed as "higher than (>)" were omitted for statistical computation purposes.

Log-transformed data were used in correlation analysis, to obtain a Gaussian-type statistical distribution (1). Parametric statistical methods were used. Analysis was initially limited to the parameters monitored by the majority of participating laboratories.

In order to verify the relationship between the most probable number (MPN) and membrane filtration culture (MF) methods in the determination of faecal coliform (FC) concentrations, different blocks of contamination levels were considered, and correlation analysis performed for each block.

3. Results and discussion

Computed correlation matrices for those parameters monitored by participating laboratories are given in Figures 1 to 5 (W = water, S = sand). In general, a highly significant correlation was found between the MPN and MF values when considering the same parameter and the same seawater sample.

Values for total coliforms (TC) and faecal coliforms (FC) showed a high level of correlation between the two techniques (MF and MPN) in the same water samples. Concentrations of faecal streptococci (FS), where this particular parameter was measured, were generally correlated to TC and/or FC levels.

Data regarding the FC content of sediments appeared to be generally correlated to data on the FC content of the corresponding water samples.

Intra-station contamination variability was observed to range up to a maximum of three orders of magnitude in some cases. A variability of one to two orders of magnitude was very frequent. This variability may be explained as mainly the result of variations in (a) pollutant dispersion phenomena and (b) meteorological patterns. A minor part of it may be due to random errors in the various measurements.

Compared to the overall variability, the differences between the MF and MPN methods were practically negligible (Figure 6).

A comparison of the data obtained from (a) slightly polluted localities and (b) heavily polluted localities indicated that in the presence of a consistent pollution source, all parameters tended to register a corresponding increase. This would account for the correlation levels given above.

The results of the correlation analysis for blocks of contamination levels are given in Appendix 1. Preliminary conclusions which may be drawn from these results are the following.

In the first place, when the contamination level is very low or practically negligible (<10 FC/100 ml) the two parameters - FC (MPN) and FC (MF) are not correlated. This may be due to the proximity to analytical thresholds and to the consequent low degree of precision, as well as to a real lack of relationship between the parameters in question in this particular range. Results could also be influenced by the limited extension of the range.

Secondly, in considering values higher than 10 FC/100 ml, a highly significant correlation emerges.

Thirdly, values lower than 100 FC/100 ml also correlate, although if the 1-10 FC/100 ml and 10-100 FC/100 ml levels are considered separately, no correlation is evident between the two subsets. Obviously, consistent variations of the contamination level may be appropriately detected in both parameters within the 1-100 FC range. Agreement between them is limited if small variations are considered.

4. Conclusions

In general, despite the remarkable intra-site variability, overall results showed that contaminated and non-contaminated sites can be appropriately distinguished and classified if a reasonable number of analytical determinations are available. The single datum is not usually significant for this purpose. An experimental design, based on an adequate assessment of intra-site variability, would be useful in the programming of seawater and sediment surveys aimed at the detection of prefixed inter-site differences.

It may be reasonably assumed that the MPN and MF methods are both adequate for distinguishing between contaminated and non-contaminated sites. For practical purposes, either method could be adopted, and selection of either method could be based mainly on operational reasons. In any case, if the random statistical fluctuations of values measured in the same sites are considered, the differences between the two methods appear to be negligible from the general point of view. The problem as to which of the two is the more reliable method may occur when particular studies are required, i.e. when the sites under examination are on the borderline from the point of view of bacteriological acceptability.

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Appendix 1

In order to verify the relationship between FC MPN and FC MF methods, different blocks of contamination levels have been considered, and correlation analysis has been carried out for each block (Figure 7). The following results have been obtained:

	Block of data	R	N	<u>P</u>
1	MPN \leq 10 and MF \leq 10	.29	41	n.s.
2	$10 \le MPN \le 100$ and $10 \le MF \le 100$.06	40	n.s.
1 + 2	MPN \leq 100 and MF \leq 100	.65	81	.001
3 + 4	MPN > 100 or MF > 100	.844	43	.001
4	MPN > 1000 or MF > 1000	.795	26	.001
2 + 3 + 4	MPN > 10 or MF > 10	.893	83	.001

Some simple preliminary conclusions may be drawn.

When the contamination level is very low or practically negligible (< 10 FC/100 ml), the two parameters, FC (MPN) and FC (MF) are not correlated.

This result may be due to the proximity to analytical thresholds and to the consequent low degree of precision, as well as to a real lack of relationship between the parameters considered, in this particular range. The limited extension of the range may also influence this result.

If values higher than 10 FC/100 ml are considered, a highly significant correlation is found.

Moreover, values lower than 100 FC/100 ml correlated too. If 1-10 FC/100 ml and 10-100 FC/100 ml intervals are considered separately, no correlation is found in the two subsets.

Evidently, consistent variations of the contamination level may be appropriately detected by both parameters in the 1-100 FC range. The agreement between them is limited, if small variations are considered.

In any case, it may be reasonably assumed that both MPN and MF methods are adequate for distinguishing contaminated and non-contaminated sites. As a rule, for practical general purposes, both methods may equally well be adopted. The selection of the MPN or MF method may be based mainly on operative reasons. In any case, if the random statistical fluctuations of values measured in the same sites are considered, the differences between the two methods appear in general negligible. The problem of which method is more reliable may be posed when particular studies are required, for instance on slightly contaminated sites.

Table 1

Results of Inter-Laboratory Exercise on Sampling and Analytical Methods

Participating institute	Testing	sam- sam- pling	sam- Total	Water	Sand and sed.	Bact. para- meters	Twa	TC	TC	TC	FC		IC FC sand sed.	ed.	FC sand s	ed.	FS sand sed.	l. Other
		point	משול הופי פ	ampres	samples		MPN	MF	MPN	MF	MPN	MF	MPN	MF	MPN	MF	MPN MF	d.
INSERM, Nice, France	14.6 - 10.8	m	18	18	1	FC, FS	1	1	+	+	+	+	1	1	ı		1	
A. Felix P.H. Lab., Tel- Aviv, Israel	25.5 - 1.8	c	32	16	16 TC,	TC, FC, FS	+	+	+	+	+	+	+	+	+	+	+	Wt°, DO, BOD ₅
Inst. Pasteur, Tunis, Tunisia	15.5 - 15.8	m	72	36	36	IC, FC	+	I	+	+	1	1	+		+	+	1	
Rudjer Bosk. Inst'., Rovinj, Yugoslavia	20.5 - 29.7	m	36	18	18 TC,	TC, FC, FS	+	+	+	+	+	+	1		+			BOD5
ISS, Rome, Italy	18.5 - 31.8		0+	24	16 TC,	TC, FC, FS	+	ŧ	+	+	I	+	+ -	1	+	!	+	At, Wt, DO, pH
Methods Arrangement prepared by ISS, April 82	1.5 - 31.7	en	8 4	24 2	24 TC,	FC, FS	+	+	+	+	+	+		+			ī	BOD5, COD, Tgg Tot.Nitr., Amm. Nitr., Org.Nitr. Tot.Phosph.

At " = air temperature (°C); Wt " \rightarrow water temperature (°C); D0 = dissolved oxygen.

TC = total coliforms; FC = faecal coliforms; FS = faecal streptococci.

Table 2 Bacteriological methods

(4 : : : : : : : : : : : : : : : : : :			Total coliforms	Lorms		Faecal coliforms	. 1			Faecal streprococci		
Institute		MPN		MF	S.	MPN	MF		MPN		W	
		Pres. test	Pres. test Conf. test		Pres. test	Conf. test	Pres. test Conf. test Pres. test Conf. test Pres. test Conf. test Pres. Test Conf. test	if, test Pres	. test C	onf. test	Pres. Test	Conf. test
S.C. 11 INSERM Nice, France	Media Time (h) Temperature (°C)				1cC. Br. 48 44	Eau pept. 7 24 44	m-FC-agar EMI 24 44	EMB/ALVBRF Az.Dx.Br. 24 48 44 44		EVA Br. 48 44	KF-agar 48 44	m-enter.agar 48 44
A. Felix Publ. Health Lab. Tel-Aviv, Israel	Media Time (h) Temperature (°C)	Lact. Br. 48±3 35±0.5	BGBB 48±3 35±0.5	m-Endo agar LES 22 35±0.5	Lact. Br. 48±3 35±0,5	EC medium 7 24 44.5±0.2 4	MF Direct MF m-FG-agar 2-; 44,5±0,2 35: 44,5±0,4 44	MF Resusc. LTB A2.Dx. 2-2 1/2 48±. 35±0.5 35±0. m-FC-agar 44.5x22 h	Br.	EVA Br. 48±3 35±0.5	MF-Direct KF-agar 48±3 35±0.5	MF-Resusc. TSB 2-2 1/2 35±0.555 KF-agar 35±0.5x48 h
Institut Pasteur de Tunis Tunisia	Media Time (h.) Temperature (°C)	McC. Br. Lact. Br. BGBB	BGBB	Endo agar (Merieux)	McC. Br. Lact. Br. BGBB 	BGBB Eau pept.	m-FC-Agar		Not inc	luded in	Not included in Tunis report	
Rudjer-Boskovic Institute Rovinj, Yugoslavia	Media Time (h) Temperature (°C)	McC. Br. 24 36±1		m-Endo agar 24 36±1	McC. Br. 24 36±1	McG. Br. 24 44.5±0.2 (gas+indole)	m-FC-Agar 24 44.5±0.2		Az.Dx.Br. B 48 36±1	BCP Az.Br. KF-Agar 48 48±3. 36±1 44,5±0.	KF-Agar 48±3 44.5±0.2	
Institute Superiore di Sanità Rome, Italy	Media Time (h) Temperature (°C)	Lact. Br. 48±3	BGBB 48±3 36±1		Lact. Br. 48±3 36±1	EC Medium 24 44.5±0.2	m-FC-Agar 24 44.5	a k			KF-Agar 48±3 44±0.2	
Analytical methods Me (arrangement Ti- proposed by Tel Istituto Superiore di Sanità in April 1982)	Media Time (b) Temperature (°C)	Lact. Br. or McC. Br. BGBB 48±3 48 36±1 36±1	BGBB 48 36±1	m-Endo agar 24 36±1	Lact. Br. or McC.Br. 48 36±1	EC Medium 24 44.5±0.2	п∽FC-Agar 24 44±0.2		Az.Dx.Br. E 48±3 36±1	EVA Br. 48 36±1	KF-Agar 48±3 44±0.2	8
Lact. Br. = Lactose Broth BGBB = Brilliant Green Lactose Bile McG. Br. = MacConkey Broth Eau pept. = Peptone water ALVBRP = Agar-lactose-Brilliant Green Phenol-Red	Broth en Lactose Bile Broth water e-Brilliant Green	Az.Dx.Br. EVA Br. = LTB = Laur BCPAz. Br. TSB = Tryp	Az.Dx.Br. = Azide Dextrose i EVA Br. = Ethyl Violet Azide LTB = Lauryl Tryptose Broth BCPAz. Br. = Bromcresol Pury TSB = Trypticase Soy Broth	Az.Dx.Br. = Azide Dextrose Broth EVA Br. = Ethyl Violet Azide Broth LTB = Lauryl Tryptose Broth MCPAz. Br. = Bromcresol Purple Azide Broth TSB = Trypticase Soy Broth	8							

Data from I.N.S.E.R.M. (France)

	T.C. _W (MPN)	T.C.W	F.C.W (MPN)	F.C.W (MF)	F.S. _W (MPN)	F.S.W (MF)	F.C.s (MPN)
T.C.(MPN)							
W							
T.C. (MF)							
W	~=						
F.C.(MPN)						_	
W		-14 -49	1	.841	.825	.832	
	-				- ×		
F.C. (MF)			.841	1	.796	.790	
F.S.(MPN)							
W		~~	.825	.796	1	.917	
F.S. (MF)		8					
W			.832	.790	.917	11	
F.C. (MPN)							
S							
	10						

N = 18

Figure 2

Data from A. Felix Public Health Laboratory (Israel)

	T.C.W	T.C.W	F.C.W (MPN)	F.C.W (MF)	F.S.W (MPN)	F.S.W (MF)	F.C.s (MPN)
T.C.(MPN) W	1	.949	.949	.947	.604	.700	.905
T.C. (MF) W	.949	1	.975	.989	.659	.719	.868
F.C.(MPN) W	.949	.975	_ 1 = 1	.971	.637	.660	.899
F.C. (MF) W	•947	.989	.971	1	.713	.739	.859
F.S.(MPN) W	.604	.659	.637	.713	1	.807	.430
F.S. (MF)	.700	.719	.660	.739	.807	1	.533
F.C. (MPN)	.905	.868	.899	.859	.430	.533	1

N = 16

 $\underline{\underline{Figure~3}}$ Correlation matrices for determining bacterial concentration

Data from Institut Pasteur de Tunis (Tunisia)

					0		
	T.C.W (MPN)	T.C.W	F.C.W	F.C. _W	F.S.W (MPN)	F.S.W (MF)	F.C.s (MPN)
r.c.(MPN) W	1		.722	.706			.241
C.C. (MF)							
F.C.(MPN) W	.722		1	.493			.340
v.c. (MF)	.706		.493	1			.042
'.S.(MPN) W						· · · · · · · · · · · · · · · · · · ·	
.s. (MF) W							40.10
r.C. (MPN) S	.241		.340	.042			1
	N = 29						

Figure 4

Data from R. Boskovic Institute (Yugoslavia)

				INDETEGE (I	O		
	T.C.W (MPN)	T.C.W	F.C. _W	F.C. _W	F.S.W (MPN)	F.S.W (MF)	F.C. _S (MPN)
c.c.(MPN)	W	2.00		2/2	0.44	0.77	0.27
W	1	.950	.977	.962	.964	.974	.804
.C. (MF)							
W	.950	1 "	.956	.986	.909	.975	.800
r.c.(MPN)	0.77	.956	1	.967	.955	.978	.811
W	.977	. 930		.707			
7.C. (MF)							
W	.962	.986	.967	1	.912	.988	.808
.S.(MPN)							
W	.964	.909	.955	.912	1	.932	.810
			-9.00		F 1 2 15		
F.S. (MF)		0.75	0.70	0.00	0.20	1	709
W	.974	.975	.978	.988	.932	11	.798
F.C. (MPN)							
S	.804	.800	.811	.808	.810	.798	1

N = 18

Correlation matrix for determining bacterial concentration

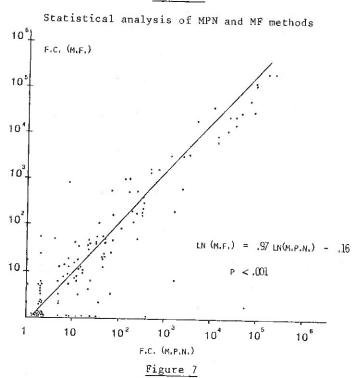
Data from Istituto Superiore di Sanità (Italy)

vata	rrom	Istituto	Superiore	d1	Sanıtà	(Italy)

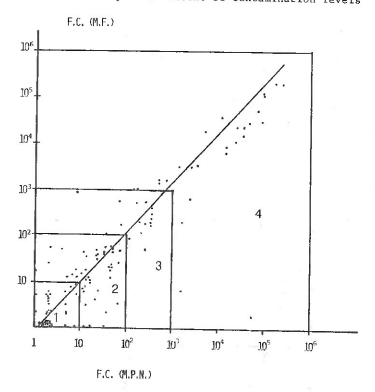
	T.C.W	T.C. _W	F.C.W (MPN)	F.C.W	F.S.W (MPN)	F.S. _W (MF)	F.C.S (MPN)
T.C.(MPN)	1		.931	.883		.425	.417
T.C. (MF)			×		i		
F.C.(MPN) W	.931		1	.892		.422	.488
F.C. (MF) W	.883	****	.892	1		.454	.406
F.S.(MPN) W)40 —			
F.S. (MF)	.425		.422	.454		1	.303
F.C. (MPN)	.417		.488	.406		.303	1

N = 25

Figure 6



Correlation analysis of blocks of contamination levels



Annex 2

AN EVALUATION OF SOME DIFFERENT METHODS FOR ENUMERATION OF FAECAL COLIFORMS FROM WATER (MOST PROBABLE NUMBER AND MEMBRANE FILTRATION TECHNIQUES)

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1. Introduction

The basic methods for the assay of pollution indicators - total coliforms (TC), faecal coliforms (FC) and faecal streptococci (FS) in water are outlined by the American Public Health Association (APHA) (1). These include the multiple tube, or most probable number (MPN) technique and the membrane filter (MF) procedure. Since the acceptance of the MF procedure for the isolation of these indicators, conflicting reports (2, 3, 4, 5) have appeared in the literature regarding the use of membrane filters as a method of evaluating the quality of water. The present study was carried out to determine the efficiency of two modified MF techniques, the LES two-step two-day procedure proposed by Stevens et al. (6) and the two-layer membrane filter procedure proposed by

Materials and methods

A total of 110 seawater samples (collected from the Alexandria area) according to the method described by APHA (1) were analysed in the present study.

The MPN procedure was performed by culturing a series of three decimal dilutions per sample, using five tubes for each dilution. Lactose Broth was used for the presumptive test, and FC medium at 44.5°C±0.5°C (water bath) was used for the confirmatory test. The MPN of FC per 100 ml was calculated from the positive EC broth using McCrady's probability tables.

In the MF procedures for FC, two duplications of each sample dilution were filtered through 0.45 μ membrane filters (MILLIPORE) for each bacterial test. One set of membranes was placed on plates of the two-layer medium (bottom layer: 37% M-FC, 1.5% agar in distilled water; top layer: 0.3% beef extract, 0.5% peptone, 0.5% lactose and 1.5% agar in distilled water). These plates were then incubated at 35°C for two hours, then at 44.5°C for 22-24 hours in watertight plastic bags in

The other membrane sets were placed on LES minimal holding agar medium (0.05% tryptose, 0.05% dextrose, 0.05% lactose, 0.05% oxgall, 0.04% sodium chloride and 1.5% agar in distilled water). These plates were incubated at 25°C for 18 hours; the membranes were then transferred to absorbent pads in tightly sealed Petri dishes that had been saturated with M-FC Broth and incubated at 44.5°C

In both MF procedures, the plates which gave 20-60 colonies per membrane were counted, and counts per 100 ml were calculated. Faecal coliform colonies were identified by their blue coloration and the crystallized deposits on their surfaces with the aid of a stereomicroscope.

Since FC densities covered many orders of magnitude, all data were expressed and analysed as logarithms (base e).

The results obtained by the different methods used for the detection of the frequency of FC are shown in Tables 1 to 3, comparing the two methods.

Table 1 shows a comparison of the frequency of FC counts/100 ml using the MPN and the LES methods. Out of 110 water samples, 14 (13%) gave FC counts of 1-99, 53 (48%) gave counts of 100-999 and 43 (39%) gave counts of 1000 or more by the MPN method. The corresponding figures for

The data presented in Table 2 show the frequency of FC counts/100 ml detected by the MPN method as compared to the two-layer method. Out of 110 water samples, 14 (13%) yielded FC counts of 1-99, 53 (48%) yielded counts of 100-999 and 43 (39%) yielded counts of 1000 or more by the MPN method. The corresponding figures for the two-layer method were 2 (1.8%), 37 (34%) and 71 (65%) respectively.

Table 3 presents a comparison of the frequency of FC counts/100 ml detected by the two-layer and the LES methods. Out of the 110 water samples 2 (1.8%) yielded FC counts of 1-99, 37 (34%) yielded counts of 100-999 and 71 (65%) yielded counts of 1000 or more by the two-layer method. The corresponding figures by the LES method were 0 (0%), 31 (28%) and 79 (72%) respectively.

The following results can be computed from these prepared tables considering the combination of the two methods. FC counts of 1-99 were found only in one sample (0.9%) by both the MPN and the two-layer method, counts of 100-999 were found in 25 (23%) by the MPN and the two-layer method, 20 for counts of 1000 or more were 43 (39%), 43 (39%) and 71 (65%) respectively.

A synthesis of the results shown in Tables 1 to 3, to show the agreement between any pair of the methods performed, is given in Table 4.

The percentage of agreement was lowest between the MPN and the LES methods (57%), followed by that between the MPN and the two-layer method (63%). The difference between these figures was not two-layer methods (91%). The difference between this figure and both the above figures was statistically significant (z=5.2 and 6 respectively, P < 0.05). Results of the statistical analysis of the data are shown in Figures 1 to 3.

Graphical presentation of the logarithms of the observed values of the LES and the two-layer methods as well as the regression line with zero intercept using the logarithmic transformation for FC counts in both methods are shown in Figure 1. This graph indicates that the fitted line is a good predictor of the relation between these two methods. The regression line with zero intercept was calculated by the method of least squares. This method fits data to an optimized line by the equation y = bx, where b is the regression coefficient or slope, y is the dependent variable, and x be: y = 0.91x in which y = FC counts/100 ml by the two-layer method and x = FC counts/100 ml by the LES method.

Comparison of the regression line with the equality line, which has a regression coefficient or slope of 1.0 and which could result if the means of the two methods were the same, shows that the regression line lies below the equality line indicating that the LES was superior to the two-layer method.

In both Figures 2 and 3 the regression lines lie below the equality lines indicating the superiority of both the two-layer and the LES methods over the MPN one.

Table 5 shows the geometric mean, arithmetic mean, standard deviation and coefficient of variation of the FC counts/100 ml as detected by the MPN, two-layer and LES methods. The highest geometric mean was found to be that of the LES method (2170) and the lowest value was obtained by the MPN method (545). The geometric mean of the two-layer method was found to be 1135. The calculated ratio of the geometric mean of FC counts using the LES method to that using the MPN method was 3.98:1, that of the two-layer method to the MPN was 2.08:1, and that of the LES method to the two-layer method was 1.9:1.

The coefficient of variation shows the degree of precision in a method of bacterial enumeration, the results with the lowest coefficient of variation being the most precise. In this study the LES method had the lowest coefficient of variation (107) whereas the MPN had the highest one (159). The coefficient of variation for the two-layer method was 140.

4. Discussion and conclusion

The results of this study revealed that the LES method enhanced FC recovery, and had a higher degree of precision than the MPN method. Higher counts were obtained in 47 of the samples examined (43%) with a geometric mean of 2170 and a coefficient of variation of 107. The MPN data reflected a geometric mean of 545 and a coefficient of variation of 159. The ratio of FC counts of the LES method to the MPN method based on the geometric mean was 3.98:1. The highest faecal coliform recovery by the LES method as compared to the MPN method in the present study might be attributed:

- (1) to the use of MILLIPORE HC brand of membrane filters, described by Green <u>et al.</u> (8) and Tobin & Dutka (9) as the most efficient one in retaining the bacteria present in the tested water;
- (2) to the use of the enriched medium (minimal holding agar) which is easily attacked and metabolized by the faecal coliforms and foster bacterial cell repair (6);
- (3) to the resuscitation period at 25°C for 18 hours that affords the injured cells the opportunity to repair themselves, multiply and divide and become insensitive to inhibitory agents in the selective media (2, 10, 11). Results obtained in the present study were in conformity with those obtained previously by several other workers including Goetzee & Pretorius (12), Public Health Laboratory Service (13) and Mara (14).

Comparable results were also obtained by Davenport et al. (15) and Green et al. (16), who attributed these to the resuscitation period (varying from 2 to 6 hours) before exposure to the elevated temperature. Other investigators reported similar results when using first an enrichment medium at low temperature (35°C for 2 to 4 hours) before exposure of the membrane to selective media at an elevated temperature of 44.5°C (4, 10, 17).

Thomas & Woodward (18) reported results disagreeing with the ones described in the present work. These authors used the standard one-step MF technique, i.e. direct incubation of the membrane on M-FC broth at 44.5°C, and observed significantly poorer recoveries of FC organisms by the MF technique than by the MPN method. They also found that the ratio of FC by MPN to MF methods was 1.3:1. This ratio was reported to be 0.92:1, 1.53:1 and 2.2:1 by Adams (19), MacCarthy et al. (20) and Slanetz et al. (17) respectively. The lower counts of the MF technique in these aforementioned studies might be attributed to the use of improper media containing substances having inhibitory effects on non-coliform Gram-negative bacilli that might also have an adverse effect on the growth of coliform organisms. Also the initial shock at 44.5°C adversely affected reproduction of metabolically injured cells.

The data obtained in the present study showed that the two-layer method was superior and more precise than the MPN method. Higher counts were obtained in 41 (37%) of the 110 positive samples with a geometric mean of 1135 and a coefficient of variation of 140; similarly, the MPN data demonstrated a geometric mean of 545 and a coefficient of variation of 159. The ratio of FC counts of the two-layer method to the MPN method based on the geometric mean was found to be 2.08:1.

In addition to the use of the MILLIPORE HC brand of membrane filters, higher FC recoveries by the two-layer method in the present study might be attributed firstly to the fact that the proposed two-layer MF procedure allows for repair and subsequent reproduction of those coliforms which have been debilitated by exposure to the aquatic environment (21); secondly, to the fact that the counts in liquid media are not as accurate as counts on solid media (14); and thirdly, to the easy and accurate counting of the characteristic blue colonies grown on the agar medium after enrichment (20, 22).

However, Stuart et al. (23) found that the two aforementioned methods gave nearly the same counts when they tested chlorinated sewage effluents with some modification in the medium. They attributed their results to the addition of glycerol and acetate, plus reducing agents to the two-layer media. Glycerol and acetate act as intermediates in the glycolytic pathway and repair the enzymatic damages of injured cells, along with reducing agents to inactivate residual chlorine.

As a result of the previous findings it was found that the MPN method gave the least counts as compared to the other two methods. The low recovery of the MPN method as compared to the LES and the two-layer methods respectively might be attributed to the false-negative results obtained in the presumptive tests due to the presence of specific soil organisms or the antagonistic action of Pseudomonas that also suppress coliform growth, so that the minimum concentration of cells required to produce visible gas in the presumptive medium is not obtained within the normal incubation time (24, 25), and the high concentration of heavy metal ions suppresses gas production by coliform bacteria (26). Also the MPN method is a biased estimator of the true density and the amount of bias depends on the number of tubes used in each dilution (27).

In this study the LES method proved to be superior and more precise in FC recovery than the two-layer method. Higher counts were found in 10 (9%) out of the 110 samples with a geometric mean of 2170 and a coefficient of variation 107. The ratio of FC counts of the LES method to the two-layer method based on the geometric mean vas found to be 1.9:1. Upon closer examination of the data, it was found that the agreement between these two methods was in 100 (91%) of the 110 samples. The great efficiency of the LES method may be attributed to the longer resuscitation period on the rich nonselective medium which contains simple carbohydrates. Although the LES method gave higher counts, it required incubation on two different media for two days to obtain the

final results. Regarding the two-layer method, it automatically shifts the culture contact from an enrichment growth substrate to the essential differential medium phase, and final results are obtained within 24 hours only. Lin (28) stated that the preenrichment incubation temperature of 35°C (used in the two-layer method) was considered the temperature of choice for the improvement of FC recovery. Ray & Speck (29) and Green et al. (16) reported that for proper enumeration of FC the resuscitation period should be completed before cell multiplication started, the time initiation of multiplication at 35°C being 4 hours. Bissonnette et al.(10) pointed out that the two-layer method avoided the limitation of enrichment techniques due to the fact that considerably more time, equipment and manpower were required for analysis. Finally, Crabow & Preez (30) proved that saturated pads were inconvenient and time-consuming, the pads tended to dry out, and the agar-based medium generally proved to yield higher counts.

In these comparisons the MPN procedure was used as the standard, but it must be noted that there are inherent shortcomings in this technique based on statistical probability, estimates of bacterial density with inherent errors (31). Membrane filtration using the two-layer agar method provides a direct accurate bacterial count, requires only 24 hours to complete, and has the advantage of saving media, chemicals, tubes, racks and space. In contrast, the MPN test requires 2 to 4 days before final results are obtained, and an excessive amount of laboratory work may be involved. In this study, from the 1600 inoculated tubes, 367 positive lactose tubes failed to confirm the presence of faecal coliforms. In other words, about 23% of all the inoculated tubes were processed through the confirmation test without any positive coliform results. One of the disadvantages of the MF technique is the high cost of membranes which have to be imported from abroad. However, Taylor & Burman (17) showed that the membrane could be washed and re-used by sterilizing in 3% (v/v) hydrochloric acid.

Following evaluation of the results obtained during the present study, it is agreed with Geldreich (22, 32, 33) that the MF technique gives the most convenient results, the LES method gives the highest FC counts and two-layer method saves time, media and labour.

5. Conclusion

Three methods were used for the detection of faecal coliforms (FC) in 110 water samples. These methods were the MPN, the LES and the two-layer MF. This work was carried out in order to determine the most efficient method for the isolation and enumeration of FC.

Comparison of the results showed the superiority and the higher degree of precision of both MF procedures over the MPN method.

It was concluded that the MF technique gives the most convenient results; the LES method gives the highest FC counts and the two-layer method saves elaborate time, media and labour.

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 $\frac{{\rm Table}\ 1}{{\rm Comparison}\ {\rm of}\ {\rm the}\ {\rm frequency}\ {\rm of}\ {\rm faecal\ coliform\ counts/100\ ml}$ detected by the MPN and LES methods

LES	MPN	1-99	FC count 100-999	1000	Total
FC count	1-99	0	0	0	0
	100-999	11	20	0	31
8 0	1000-	3	33	43	79
	Total	14	53	43	110

Table 2

Comparison of the frequency of faecal coliform counts/100 ml detected by the MPN and two-layer methods

			FC count		
Two-layer	MPN	1-99	100-999	1000	Total
FC count	1-99	1	1	0	2
	100-999	12	25	0	37
	1000-	1	27	43	71
	Total	14	53	43	110

Table 3

Comparison of the frequency of FC counts/100 ml detected by the two-layer and the LES methods

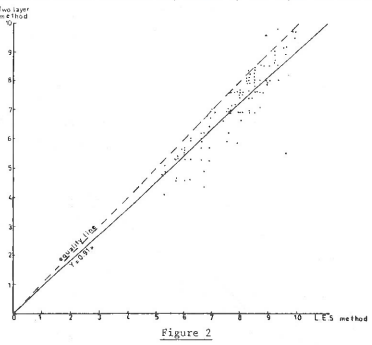
	Two-layer		FC count		
LES		1-99	100-999	1000	Tota
C count	1-99	0	0	0	0
	100-999	2	29	0	31
	1000-	0	8	71	79
	Total	2	.37	71	110

 $\frac{\text{Table 4}}{\text{Agreement between the MPN, the two-layer and the LES methods}}$ in the estimation of faecal coliforms

First method			FC co	FC counts			
	Second method	Equal number by the two methods	Higher counts by 1st method than 2nd method	Lower counts by 1st method than 2nd method	Percentage of agreement		
MPN	LES	63	0	47	57		
MPN	Two-layer	69	1	40	63		
LES	Two-layer	100	10	0	91		

Figure 1

Comparison of faecal coliform log counts by two-layer and LES methods



Comparison of faecal coliform log counts by MPN and two-layer methods

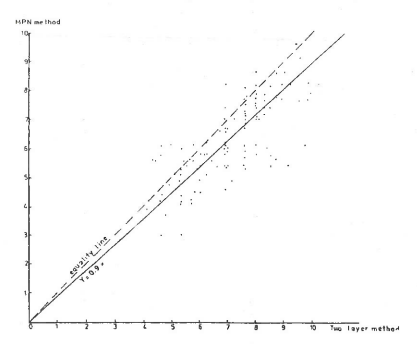
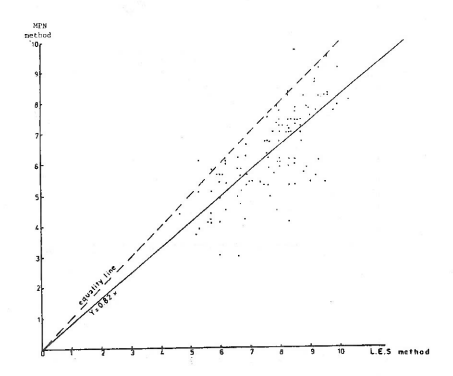


Figure 3

Comparison of faecal coliform log counts by MPN and LES methods



Annexe 3

ETUDE COMPARATIVE DES METHODES UTILISEES EN NUMERATION BACTERIENNE DES EAUX COTIERES DE LA MEDITERRANEE

par
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1. Introduction

La présente étude, entrant dans le cadre des travaux du Service Commun N°11 de l'Institut national de la Santé et de la Recherche médicale (INSERM) pour le Programme MED POL Phase II, a eu pour but de comparer deux méthodologies de numération des germes fécaux (colibacilles et streptocoques) en eau de mer des zones côtières.

A Nice, trois zones ont été choisies :

- la plage de Coco Beach, formée de petites criques rocheuses, située près du Cap de Nice;
- la plage du Forum, au centre de la grande plage de galets de la Baie des Anges, au débouché des principales artères de la Ville de Nice;
- le port de commerce de Nice (Port Lympia), implanté à l'est de la cité; ce site a été sélectionné en considération de sa forte charge bactérienne présumée.

L'exercice d'essaí a été conduit de la mi-juin à la mi-août 1982 au cours de six opérations de prélèvements d'eaux de mer côtières effectuées en chacun des trois lieux précités (voir figure 1), soit au total 18 prélèvements.

2. Matériel et méthodes

Prélèvements d'eau de mer

Les prélèvements ont été effectués avec des flacons en verre stérilisés de l litre, aux distances suivantes :

Coco Beach :

1 m du bord (rocher)

Forum :

3 m du bord (galets)

Port de Commerce :

0,50 m du bord (béton).

Pour chacun de ces trois sites, 1 à 2 litres ont été échantillonnés à chaque fois au même endroit, à 20 cm sous la surface de l'eau, et les analyses ont commencé une heure après au plus tard.

Volumes d'eau étudiés

Lieux à Nice	filtre	es (ml) És sur m néthode	embrane	Volumes (ml) d'eau introduits en tubes (méthode MPN)		
Plage de Coco Beach	100	10	1	10	1	1 à 10 ⁻¹
Plage du Forum	100	10	1	10	1	1 à 10 ⁻¹
Port de Commerce	100	1	1 à 10 ⁻¹	1	1 à 10 ⁻¹	$1 \text{ à } 10^{-2}$

Filtration par membranes (MF)

Les membranes utilisées (SARTORIUS) étaient de 47 mm, 0,45 u de porosité.

Chaque volume d'eau et chaque dilution ont été filtrés 5 fois pour culture sur gélose spécifique à l'aide d'une rampe de filtration (MILLIPORE).

Tubes multiples (MPN)

Chaque volume et chaque dilution ont été répartis dans 5 tubes de verre avec bouchons en coton entouré de gaze.

Prises des échantillons

Elles ont été faites avec des pipettes automatiques PIPETMAN 5000 μl et 1000 μl avec des cônes stérilisés.

Géloses utilisées

Coliformes fécaux

- a) m FC Broth base (DIFCO) additionné d'acide rosolique
- b) EMB (PASTEUR)
 Gélose Lactosée Vert Brillant Rouge de Phénol (PASTEUR).

Streptocoques fécaux

- a) KF-streptococcus-agar (DIFCO) additionné de TTC
- b) Enterococcus-agar (DIFCO).

Bouillons utilisés

Coliformes fécaux

- a) Mac Conkey Broth (DIFCO) avec cloche
- b) eau peptonée exempte d'indole (PASTEUR).

(2) Streptocoques fécaux

- a) Azide-Dextrose Broth (DIFCO)
- b) EVA Broth (DIFCO).

Incubation

L'incubation a été de 48 h pour les géloses et les bouillons à streptocoques fécaux, ainsi que pour les bouillons Mac Conkey.

Pour les géloses FC, EMB, au Vert Brillant-Rouge de Phénol et pour l'eau peptonée, l'incubation a été arrêtée à 24 h.

Toutes les cultures ont été faites à 44 °C.

Lecture des résultats (effectués par la même personne)

Coliformes fécaux

- a) Filtrations sur membranes :

 colonies bleu ciel à bleu foncé; pour confirmation certaines colonies ont été repiquées sur gélose EMB ou gélose Lactosée au Vert Brillant-Rouge de Phénol.
- b) Tubes multiples :
 - croissance, virage du lactose, 5-7 mm de gaz
 - en Mac Conkey, confirmation de tous les positifs en eau peptonée.

Streptocoques fécaux

- a) Filtrations sur membranes :
 - colonies roses à rouge foncé sur gélose KF
 - repiquage éventuel sur gélose Enterococcus.
- b) Tubes multiples :
 - croissance et culot foncé en bouillon Azide
 - confirmation de tous les tubes positifs en bouillon EVA.

Résultats

La présente étude ayant pour but de comparer deux méthodes d'analyse, la moyenne de chaque mesure est présentée avec un intervalle de confiance à 95%. De manière à visualiser les résultats, ces derniers ont été regroupés dans les tableaux 1 et 2 et les figures 2 à 7 (échelle semi-logarithmique, base 10 pour les figures).

3.1 Coliformes Fécaux

Les résultats des analyses (figures 2, 4 et 6) par les deux méthodes (filtration sur membranes et tubes multiples) sont compatibles entre eux.

La seule exception concerne le prélèvement effectué à la plage du Forum, le 26 juillet, où un nombre élevé de coliformes fécaux a été mis en évidence (FM = moyenne de 800; tubes multiples = moyenne de 8). Une explication possible de cet écart important serait la présence de nombreux germes alkalinisants (100 colonies rouges par 100 ml sur FM en gélose FC) qui auraient faussé la lecture des bouillons Mac Conkey (tubes multiples).

3.2 Streptocoques fécaux

Les résultats obtenus (figures 3, 5 et 7) ne sont guère différents d'une méthode à l'autre. Pour les deux groupes de germes fécaux, l'intervalle de confiance est plus grand pour la méthode des tubes multiples (MPN).

4. Conclusion

Il apparaît que les deux méthodes de numération - filtrations sur membranes (MF) et tubes multiples (MPN) - donnent des résultats similaires, à l'exception du cas spécial des eaux portuaires.

Ces résultats restent à être interprétés en fonction des données recueillies par l'ensemble des laboratoires ayant participé à l'étude.

Numérations des coliformes fécaux pour 100 ml d'eau de mer (moyennes et intervalles de confiance à 95%)

New York	D-A-	Filtration sur membranes Nombre le plus probat (5 membranes) (5 tubes)					obable	le	
	Date	moyenne	lim. inf.	lim. sup.	moyenne	lim. inf.	lim.	sup.	
	14.06.82	4	1	6	2	0,5	7		
	21.06.82	9	6	12	17	5	46		
Plage	02.07.82	2	1	5	37	23	41		
de	09.07.82	25	18	31	70	23	170		
oco Beach	26.07.82	14	10	18	7	1	17		
	03.08.82	8	6	10	12	3	28		
			,			8		3 /	
	14.06.82	165	120	210	79	25	190		
	21.06.82	2	1	2	2	1	2		
Plage	02.07.82	3	1	5	2	1	7		
du	09.07.82	13	10	15	17	5	46		
Forum	26.07.82	800	660	940	8	1	19		
	03.08.82	6	4	8	5	0	13		
			 						
	14.06.82	600	480	720	2400	2400	> 2400		
	21.06.82	1340	830	1850	490	170	1300		
Port	02.07.82	480	220	740	170	50	460		
de	09.07.82	3020	2110	3930	2400	680	7500		
Commerce	26.07.82	1600	1340	1860	720	560	880		
	03.08.82	3120	2470	3770	1300	350	3000		

Tableau 2

Numérations des streptocoques fécaux pour 100 ml d'eau de mer (moyennes et intervalles de confiance à 95%)

	Date	Filtra	tion sur me (5 membrar		Nombre le plus probable (5 tubes)		
		moyenne	to the teacher	lim. sup.	moyenne	lim. inf.	lim. sup.
	14.06.82	51	35	67	23	7	70
	21.06.82	25	11	32	8	1	19
Plage	02.07.82	10	4	16	7	1	17
de	09.07.82	37	23	41	79	25	190
Coco Beach	26.07.82	12	9	15	8	1	19
	03.08.82	1	0	2	4	0	11
				ACA	<u> </u>	1.00	11.5
	14.06.82	85	74	97	27	9	80
	21.06.82	22	11	32	2	1	7
Plage	02.07.82	21	13	28	46	16	120
du	09.07.82	13	10	16	14	4	34
Forum	26.07.82	9	5	13	13	3	31
	03.08.82	2	1	3	2	0	7
					, 根 使		<i>p</i>
	14.06.82	1800	1520	2080	350	120	1000
	21.06.82	3870	2620	5120	1100	310	2500
Port	02.07.82	680	350	1010	230	70	700
de	09.07.82	2400	1910	2890	1700	430	4900
Commerce	26.07.82	720	560	880	110	20	250
	03.08.82	1280	1020	1540	230	70	700

<u>Figure 1</u>

Zones de prelevements à Nice

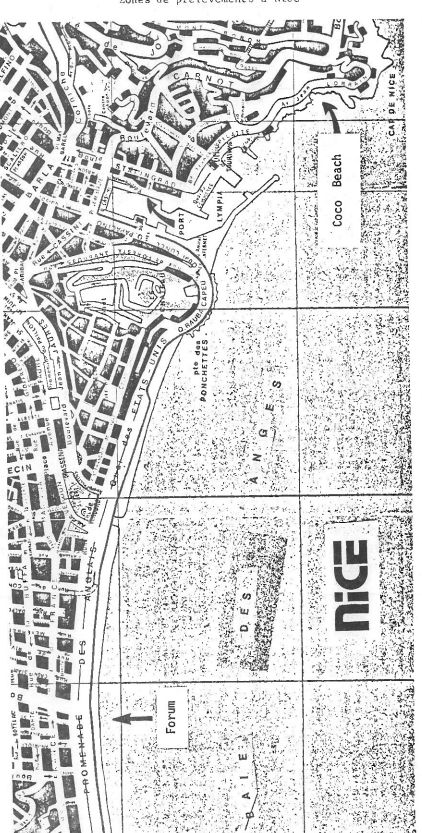


Figure 2

Intervalles de confiance à 95% pour le taux de Coliformes fécaux (plage de Coco Beach, 2 méthodes de numération)

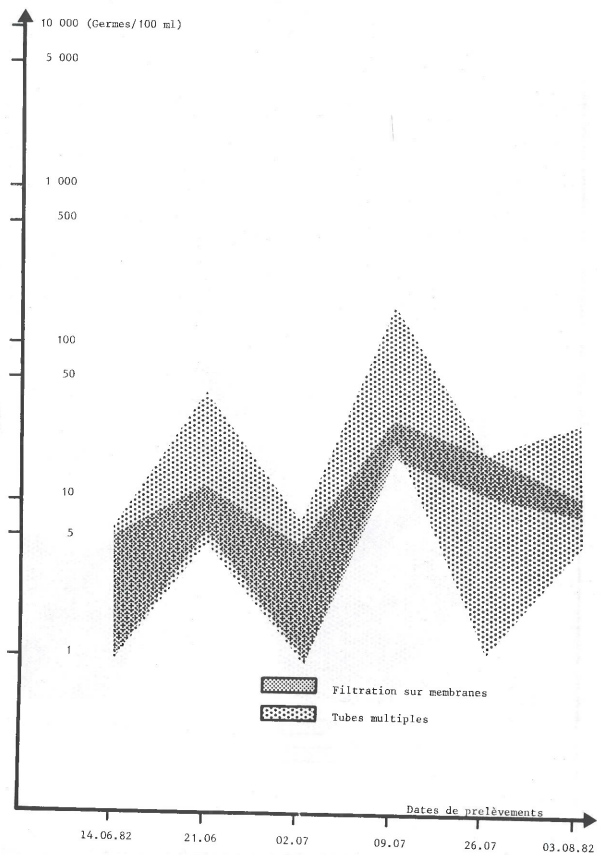


Figure 3

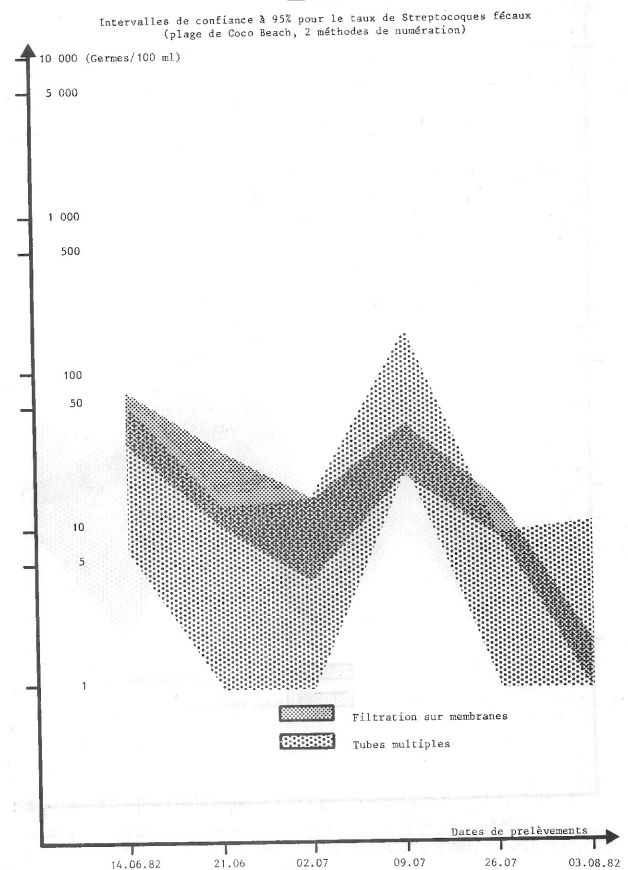


Figure 4

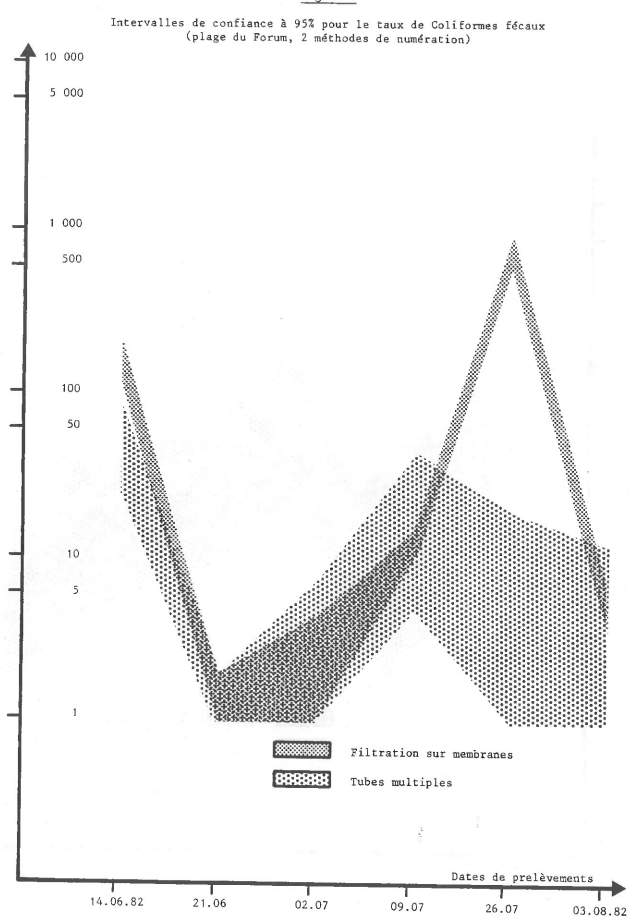


Figure 5

Intervalles de confiance à 95% pour le taux de Streptocoques fécaux (plage du Forum, 2 méthodes de numération)

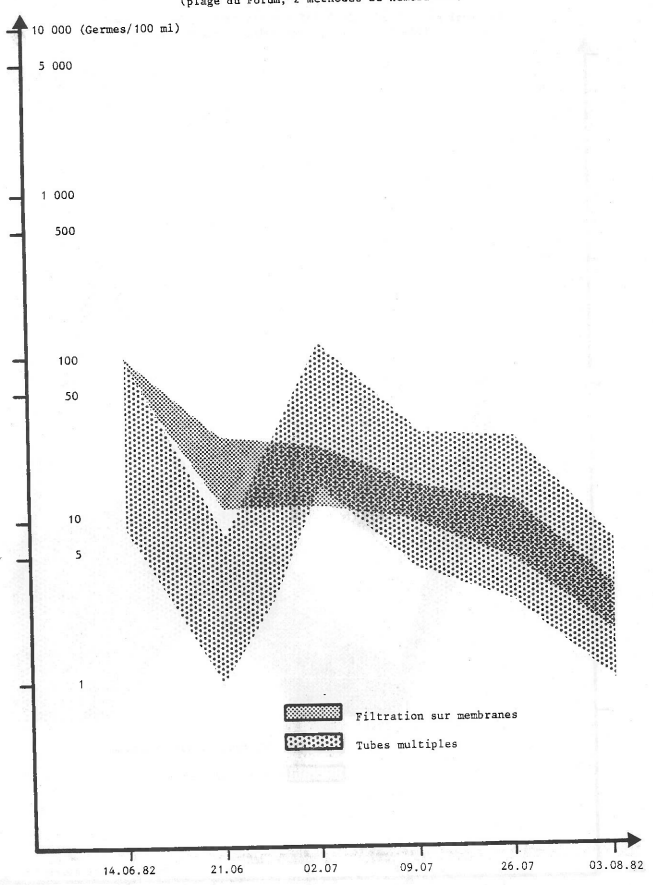
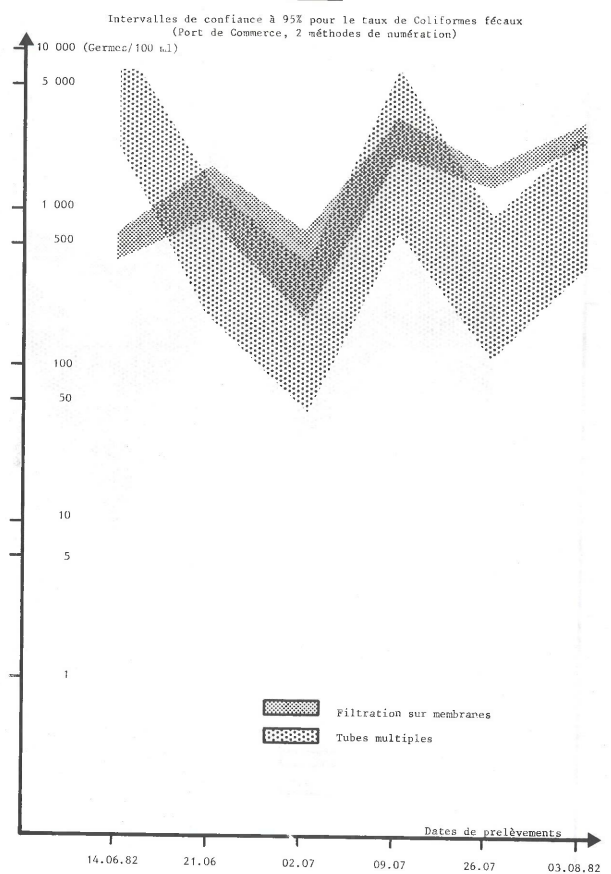
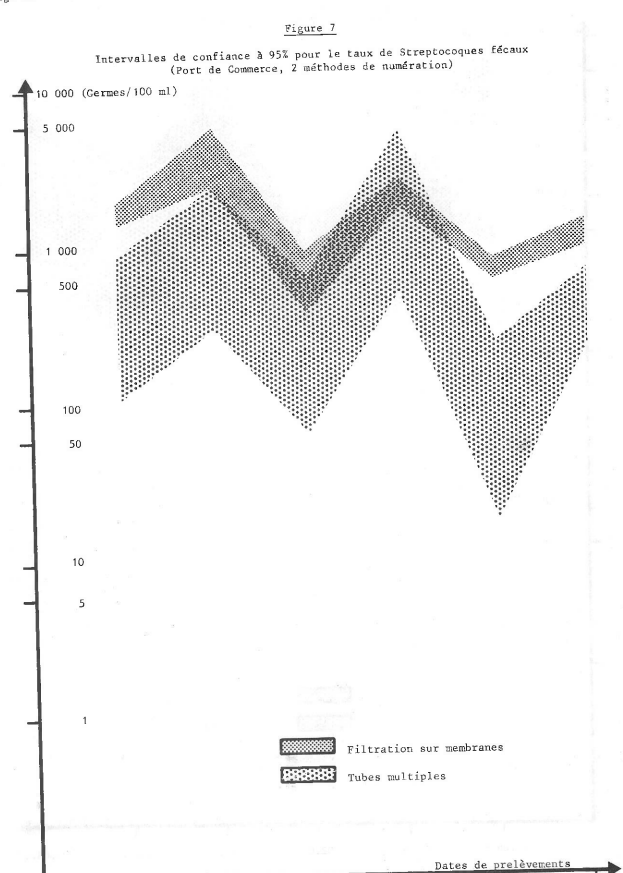


Figure 6





03.08.82

09.07

02.07

14.06.82

21.06

26.07

Annex 4

COMPARISON OF METHODS FOR MONITORING INDICATOR ORGANISMS IN MARINE WATER

by
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1. Introduction

The purpose of the present study was to compare the most probable number (MPN) and membrane filtration (MF) methods in monitoring the major bacteriological parameters for assessing the quality of coastal water in the Mediterranean. This would be useful in the eventual recommendation of the most suitable one as a standard procedure.

In previous comparisons of these two methods in coastal water (1, 2), the number of coliforms obtained by the MPN procedure was usually higher than that obtained by the MF method, which raised the question of the rate of recovery of stressed organisms by the accepted one-step MF method. An attempt was made in the present study to elucidate this point by examining parallel samples for faecal coliforms and faecal streptococci with and without resuscitation.

2. Materials and methods

2.1 Testing period

Sampling started on 25 May 1982 and continued until the first week in August. Two of the sampling points selected on the first sampling day gave very low results and were replaced by point and only five in the other two.

2.2 Sampling points

No means of sampling at the pollution source itself, which was approximately 300 m offshore, was available. The closest accessible location was approximately 500 m north of the sewage outfall at Tel-Baruch, where bathing is forbidden. The other sampling points were: Sheraton Beach, circa 900 m south of the outfall, and Bugrashov Beach, circa 1800 m south of the outfall.

Two litres of water and about 200 g of sediment were taken at each point and transported to the laboratory within $1-1\ 1/2$ hours. The water temperature was measured at the time of sampling.

Total coliforms

- MPN: Presumptive test in Lactose Broth confirmed in Brilliant Green Bile Broth, both incubated at 35°C±0.5°C for 48±3 hours. Four to five dilutions of each sample were inoculated in a 5-tube series and the appropriate three dilutions were selected for reading results.
- MF: Two to three dilutions were filtered from each sample. The membranes were placed on M-endoagar LES and incubated at $35^{\circ}\text{C}\pm0.5^{\circ}\text{C}$ for 22 hours.

Faecal coliforms

- MPN: From each positive tube of the presumptive test for total coliforms two drops were inoculated into a tube of EC medium using a Pasteur pipette. The tube was incubated at 44.5°C±0.2°C (water bath) for 24 hours. Tubes showing any amount of gas were considered positive.
- MF: A double series was filtered from each dilution. One was placed directly on M-FC-agar and incubated at 44.5°C±0.2°C, the other was placed on pads saturated with Lauryl Tryptose Broth (3) (in the covers of the Petri dishes) and incubated at 35°C±0.5°C for 2 2 1/2 hours, then transferred to the M-FC-agar and incubated at 44.5°C±0.2°C for circa 22 hours.

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Faecal streptococci

- MPN: Presumptive test in Azide-Dextrose Broth, as outlined for total coliforms, confirmed in EVA Broth, both incubated at 35°C±0.5°C for 48±3 hours.
- MF: In view of the existence of two sets of directions for faecal streptococci, one calling for incubation at 35°C±0.5°C (3) and the other at 44°C±0.2°C (4), both temperatures were tested on one sampling day. Counts obtained at 35°C±0.5°C were considerably higher, especially in the least polluted points, therefore all further examinations were incubated at 35°C±0.5°C for 48±3 hours.

A double series was filtered for each dilution, one set of filters was placed directly on KF-agar and the other was resuscitated on pads saturated with Trypticase Soy Broth for $2-2\ 1/2$ hours at $35^{\circ}\text{C}\pm0.5^{\circ}\text{C}$ then transferred to KF-agar and incubated for 2 days.

From each filter, several colonies representative of the various groups were selected and transferred to blood agar plates, then to brain heart infusion broth. They were examined microscopically (Gram stained) and for production of oxidase.

2.3 Examination of sediment

From the sediment samples, 50 or 100 g portions (wet weight) were weighed and 10 volumes of phosphate buffer with 0.1% peptone added to each. The diluted samples were shaken vigorously by hand and as soon as most of the sand had settled the liquid was decanted and used for inoculation. The results obtained were multiplied by 10 and calculated for 100 g of sample.

2.4 Other parameters

Dissolved oxygen and BOD_5 were determined in all water samples. Salinity was measured only on one sampling day.

2.5 Estimation of precision

The precision of the membrane filtration technique was checked by filtering triplicate samples from a triplicate dilution series of faecal coliforms and faecal streptococci in marine water (using strains isolated during the present work), and subjecting the results to analysis of variance to determine the standard error. The number of organisms in the suspensions was also determined on plate count agar.

Results

The data obtained for each sampling point are presented in Tables 1 to 3. Results show that when total coliforms were chosen as the parameters for assessing the bacterioloigcal quality of water or sediment, and the limit set at 1000 organisms per 100 ml of water or 100 g of sediment, one sample of water and two samples of sediment from the Sheraton Beach would exceed that limit when monitored by the MPN method, while complying with it by the MF procedure. In the case of faecal coliforms, using the MPN method, this limit was exceeded only in one sample of sediment. All other samples fell into the same classification by either method. The number of faecal streptococci recovered was much higher by the MF method with KF-medium and the resuscitation step increased their number considerably. The incubation temperature of 35°C±0.5°C was found to be more efficient in recovering faecal streptococci (as illustrated in Table 4).

Salinity was measured on one day only (18 July). Levels were $35.2^{\rm O}/{\rm oo}$ at Tel-Baruch and $35.49^{\rm O}/{\rm oo}$ at the Sheraton and Bugrashov Beaches.

The estimation of precision is summarized in Table 5.

4. Discussion

The results for total coliforms were higher by the MPN procedure in 75% of the samples, by the MF method in 15% of the samples and equal for both methods in 10% of the samples. When all data were subjected to the Student's t-test, the difference was not statistically significant. However, when the two unusually high readings at Tel-Baruch on 18 July were excluded the difference was statistically significant.

Faecal coliform readings were higher by the MPN technique in 44% of the samples, by the MF in 31% of the samples and equal by both methods in 25% of the samples. Differences were not statistically significant.

Several authors have reported the MPN procedure as being more efficient in recovering total and faecal coliforms from stream water and sewage (5) from wastewater effluent (6) and from seawater (7). On the other hand, Dutka & Tobin (8) achieved maximum estimates of coliforms in the Lower Great Lakes region of Canada by the MF procedure with M-Endo-agar LES. They compared four procedures and found that each was selective for different genera of Enterobacteriacea.

According to Rose et al. (9), a two-layered medium proposed by them, which included two hours' pre-incubation at 35°C, recovered 3.8 to 7 times more faecal coliform colonies from marine water than the direct M-FC procedure. In the present study, the additional resuscitation step recovered a larger number of faecal coliforms only in a few samples and its use would not therefore appear to be justified, particularly in clean beaches, in view of the considerable amount of time and work involved.

In polluted water however (Tel-Baruch), the difference between the number of faecal coliforms recovered after resuscitation and without it was statistically significant (P 0.1).

Since the data presented here were collected during a short period of time in a limited area, it appears that for final selection of the most suitable method for monitoring faecal coliforms in the Mediterranean more data from a wider area should be compared.

Counts of faecal streptococci were considerably higher at an incubation temperature of $35^{\circ}\text{C}\pm0.5^{\circ}\text{C}$ than at $44.5^{\circ}\text{C}\pm0.2^{\circ}\text{C}$ (Table 4) and increased significantly with a 2 - 2 1/2 hour resuscitation period on Trypticase Soy Broth.

In 90% of the samples, counts were much higher by the direct MF method at $35^{\circ}C\pm0.5^{\circ}C$ than by the MPN procedure (Tables 1 to 3) and analysis of all data showed that the difference was statistically significant (P 0.1). After resuscitation the difference was statistically significant at a higher level (P 0.01).

No explanation can be offered for the enormous discrepancy between the recovery rate of faecal streptococci by MF on KF-medium and MPN methods. Among the colonies that were confirmed, 10-30% were not streptococci (there were micrococci and even a few strains of Pseudomonas) and this rate did not account for the difference. The ratio between the MF and the MPN densities was from 0.6 to 92, with 220 in one sample and 355 in another. Kenner et al. (10), who introduced the KF-medium, obtained ratios of 0.61 to 20 with one of 200. Different media and procedures for recovering faecal streptococci from water have been compared by various authors (10-14) and all of them reported that KF and PSE-agar gave the highest recovery rates. Several authors prefer the PSE-medium, as it requires only 24 hours incubation versus 48 hours on KF-medium. Daust & Litzky (12) obtained the highest counts from marine water on PSE agar and from sewage on KF-agar; Brodsky & Schiemann (13) found that KF medium was not as selective as PSE agar - only 65% of all colonies they confirmed were faecal streptococci compared to 90% from PSE.

In conclusion it can be said that the data presented in this study, as well as that available in the literature, indicate that for faecal streptococci the MF procedure with KF-medium is superior to the MPN method, as it gives better results in a shorter time (two days instead of four) with less work involved. Since the PSE-medium has the advantage of shorter incubation time, and has been reported by some authors to be more selective in some environments, it may be desirable to compare the two media for selecting the most suitable one for Mediterranean waters.

Acknowledgement

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Table 1

Comparison of indicator organism densities per 100 ml of water or 100 g of sediment at the Tel-Baruch coast by MPN and MF methods

Dare	Temp.	Total	Total coliforms	Fa	Faecal coliforms	18	Fae	Faecal streptococci	occi		
	ວຸ	MPN	MF-D	MPN	MF-D	MF-R	MPN	MF-D	MF-R	00	BOD5
Water								55			
6.6.82	25	920	340	350	170	011	7.6	0.07.00	0000	(,
22.6.82	27	35000	5600	4900	8000	22000	27,00	2500	3700	ກ ເ	00 i
4.7.82	27.8	28000	24000	22000	11000	22000	0044	2000	3500	Λ. Ο ι	2.0
18.7.82	28.5	35000	100000	11000	38000	41000	0000	1,000	2000	ر ر ، ر	5.15
1.8.82	29.5	160000	95000	92000	30000	48000	11000	10000	21000	5.2	6.2
MEAN*		22000	13000	8200	7000	10000	2000	7000	11000		
SD		7	11	00	6	13	12	2	2		
Seliment											
6.6.82		17000	7500	3500	7600	0027	24000	71000	000		
22.6.82		4600	2800	0065	4400	20000	16000	21000	22000		
4.7.82		2400	15000	3500	0009	3000	24000	15000	22000		
8.7.82		16000	200000	9200	14000	14000	2200	2000	38000		
1.8.82		17000	2800	17000	006	800	3300	8100	7800		
MEAN*		10000	28000	9009	4400	5000	9200	13000	28000		
SD		2	14	2	6	4	3	7	2,800,0		

* Geometric mean rounded up to 2 significant figures MF-D: membrane filtration, direct MF-R: membrane filtration with resuscitation

Table 2

Comparison of indicator organism densities per 100 ml of water or 100 g of sediment at the Sheraton Beach by MPN and MF methods

	Temp.	Total	Total colitorms	E C	Faecal coliforms		Faecal	al streptococci	cci	;	
Date	ູ້ວ	MPN	MF-D	MPN	MF-D	MF-R	MPN	MF-D	MF-R	00	BODS
Water											
25.5.82	25	240	160	240	50	270	42	34	ı	7.3	3.1
6.6.82	25	1600	320	220	110	110	. 2	710	1800	6.5	1.5
22.6.82	2.7	130	14	00	80	42	2	044	1200	5.5	1.2
4.7.82	27.8	110	110	26	40	42	70	3500	7000	0.9	1.6
18.7.82	28.5	23	26	8	18	10	170	1100	2700	0.9	0.9
1.8.82	29.5	64	19	13	9	9	170	076	2500	5.5	9.0
MEAN*		190	09	32	24	38	14	580	2500		
SD	20	4	4	5	en .	4	11	ıc	2		
Sediment					R V	a a	8	a -	- 20		
25.5.82		24000	096	1300	1200	1200	310	820			
6.6.82		9400	390	1100	006	1200	4600	7500	18000		
22.6.82		490	20	50	20	40	067	3200	17000		
4.7.82		50	04	20	50	<20	330	7000	0006		
18.7.82		67	50'	67	70	20	1100	2000	17000		
1.8.82		20	10	< 20	20	<20	13	42	420		
MEAN∻	V E	390	73	76	110	70	390	1800	7100		
SD		18	9	∞	9	10	7	7	5		

* Geometric mean rounded up to 2 significant figures MF-D: membrane filtration, direct ME-R: membrane filtration with resuscitation

Table 3

Comparison of indicator organism densities per 100 ml of water or 100 g of sediment at the Bugrashov Beach by MPN and MF methods

$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Date	Temp.	Tota	Total coliforms	ø	NGX.	Faecal coliforms	iforms		N	Faecal streptococci	cocci		
6.82 25 49 2 2 2 2 2 2 2 350 970 710 1700 6.5 5.7 5.8 2 27.8 49 8 2 2 2 36 49 1300 150 150 5.7 5.8 2 27.8 170 100 110 38 60 49 1300 150 5.7 5.8 2 29.5 33 40 23 39 34 220 400 1000 5.7 5.7 5.2 29.5 130 200 50 5.7 5 5 6 7 7 5 5 144 50 1000 5.7 5 7 5 5 6 7 7 5 5 144 50 1000 5.7 5 7 5 5 6 7 7 5 5 7 144 50 1000 12000	Water					MEN	MF	Q	MF-R	MPN	MF-D	MF-R	8	BOD ₅
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	6.6.82 22.6.82 4.7.82 18.7.82 1.8.82	25 27 27.8 58.5 29.5	49 49 170 46 33	100	0 2 0 8 5	2 2 110 46 46 23	36	2 2 8 2 2	350 6 60 19 34	970 2 49 130 220	710 150 1300 410 400	1700 150 3500 1100 1000	6.5 5.7 6.3 6.0	0.6 1.5 1.0 0.6
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	MEAN* SD		57	18		14	8		12 5	44	500	1400		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	diment						2							
100 42 37 18 24 330 2300 5 4 6 3 3 3 3 3	6.6.82 2.6.82 4.7.82 8.7.82 1.8.82		230 130 700 20 50	200 20 160 20 20 20 10		50 20 700 20 20	50 20 100 100 10		200 20 20 10 20	80 490 1400 220 330	3600 2000 8000 1600 650	6100 12000 24000 1400 1300		
	IN*		100	45		37 6	18		24	330	2300	5000		

* Geometric mean rounded up to 2 significant figures MF-D: membrane filtration, direct MF-R: membrane filtration with resuscitation

Table 4 Effect of incubation temperature on recovery of faecal streptococci on KF-medium (number of organisms per 100 ml water or 100 g sediment)

			Incuba	tion temperature		
Location			35°C±0.5°C	44	.5°C±0.2°C	
		MF-D	MF-R	MF-D	MF-R	- 1.15 0.15
Tel-Baruch	water	3600	3500	2300	2400	
	sediment	21000	27000	22000	20000	
Sheraton	water	440	1200	400	20000	
	sediment	3200	17000	900	4800	
Bugrashov	water	150	930	40	170	
	sediment	2000	12000	200	8800	
			1. * 1	8 8		700
MEAN*		1800	5300	470	2300	
SD		6	4	13	6	
						800

Geometric mean rounded up to 2 significant figures

MF-D: membrane filtration, direct
MF-R: membrane filtration with resuscitation

Table 5 Estimation of Precision of Membrane Filtration

			1 74.634			
Total c	oliforms	Faecal o	coliforms	Faecal st	reptococci	
Mean	SE	Mean	SE	Mean	SE	61 411
132 143	2.3 7.3	119 123	3.2 11.7	15 22	0.33	
131	4.0	110	13.1	22	5.5	
	8.6	= 1 × ×	17.8		6.5	
	NS		NS		NS	
	Mean 132 143	132 2.3 143 7.3 131 4.0	Mean SE Mean 132 2.3 119 143 7.3 123 131 4.0 110	Mean SE Mean SE 132 2.3 119 3.2 143 7.3 123 11.7 131 4.0 110 13.1 8.6 17.8	Mean SE Mean SE Mean 132 2.3 119 3.2 15 143 7.3 123 11.7 22 131 4.0 110 13.1 22 8.6 17.8	Mean SE Mean SE Mean SE 132 2.3 119 3.2 15 0.33 143 7.3 123 11.7 22 3.3 131 4.0 110 13.1 22 5.5 8.6 17.8 6.5

Annex 5

A COMPARISON OF MICROBIOLOGICAL METHODS FOR MONITORING INDICATOR ORGANISMS IN SEAWATER AND SAND

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1. Introduction

The purpose of this study was to compare two different analytical methods, most probable number (MPN) and membrane filtration (MF) for determination of the faecal coliform density in coastal seawater samples. In addition, total coliforms and faecal streptococci were determined in water samples and total and faecal coliforms and faecal streptococci in sand samples.

The following data were also determined: sea and sky conditions, surface current, wind, last relevant precipitation, air and water temperature, dissolved oxygen (DO) and pH.

2. Materials and methods

2.1 Testing periods

Sampling was performed between 18 May and 31 August 1982, the following being the sampling dates: 18.5.82; 1.6.82; 15.6.82; 26.6.82; 13.7.82; 27.7.82; 10.8.82; 31.8.82.

Hydrographic and meteorological conditions on each sampling day are listed in Table 1.

2.2 Sampling points

The following sampling stations (on the coast of the Province of Latina, between Torre Astura and Torre Paola) were selected (Figures 1 and 2):

- No. 1: "Canale Caterattino" a small canal receiving sewage discharges only in the summer. Water samples were collected near the outfall of the canal into the sea. Sand samples were collected on the shoreline, near the outfall of this canal.
- No. 2: beach, at km 25.6 on the coast road clean area. Samples of seawater and sand were collected.
- No. 3: eliminated.
- No. 4: "Rio Martino" canal receiving most of the municipal industrial wastewater of Latina (80 000 inhabitants approx.). The samples were collected from the bridge, at km 17.7 on the coast road.
- No. 5: at km 17.1 on the coast road (approx. 500 m north-north west of the outfall of Rio Martino into the sea) under the possible influence of this pollution source.
- No. 6: beach, located 500 m south east of station No. 7.
- No. 7: "Fosso Mascarello" canal receiving the municipal and industrial wastewater of the area (40 000 inhabitants approx.).
- $\underline{\text{No. 8}}$: beach, located 500 m north west of station No. 7.

2.3 Sampling methods

Water samples were collected in sterile 1 litre bottles. In sampling points 2, 5, 6 and 8 (seawater) the samples were collected by hand at about 10 m from the shoreline. In sampling points 1, 4 and 7 collection was performed by sampling device. In sampling points 1, 2, 5, 6 and 8 samples were collected using a sterile wide-mouthed container.

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All samples were collected between 9 a.m. and 3 p.m. and transported in cooling containers $(4-10^{\circ}\text{C})$ protected against ultraviolet irradiation and examined in the laboratory. The maximum transportation and storage time was 24 hours).

In all, 56 water and 40 sand samples were collected and analysed.

2.4 Analytical methods

2.4.1 Water samples

Total coliforms (MPN): presumptive test tubes of Lactose Broth were incubated at 36°C±1°C (total incubation time = 48 hours). Positive tubes were confirmed in Brilliant Green Lactose Bile Broth tubes and incubated at 36°C±1°C for 48 hours.

Faecal coliforms (MPN): positive test tubes of Lactose Broth were confirmed in EC Broth tubes and incubated at 44.5 °C±0.2 °C for 24 hours.

Faecal coliforms (MF): each sample (or its dilution) was filtered by MF. The membrane was then incubated on M-FC-agar at 44.5°C±0.2°C for 24 hours.

Faecal streptococci (MF): each sample (or its dilution) was filtered by MF. The membrane was then incubated on KF-agar at 44°C±0.2°C for 48±3 hours.

Other parameters (dissolved oxygen, pH, air and water temperature) were determined by standard technical methods.

2.4.2 Sand samples

Total coliforms, faecal coliforms and faecal streptococci were determined also in sand samples collected from the shoreline. Sand was examined by adding approximately 50 g of sand to a beaker containing 50 ml of physiological saline, followed by analysis for total coliforms, faecal coliforms and faecal streptococci, as follows:

Total coliforms (MPN): presumptive test tubes of Lactose Broth were incubated at 36°C±1°C. Positive tubes were confirmed in Brilliant Green Lactose Bile Broth tubes and incubated at 36°C±1°C for 48 hours.

Faecal coliforms (MPN): positive test tubes of Lactose Broth (see total coliforms) were confirmed in EC broth tubes and incubated at 44.5°C±0.2°C for 24 hours.

Faecal streptococci (MF): the sample was filtered by MF and the membrane incubated on KF-agar at 44°C±0.2°C for 48±3 hours.

Results

Results obtained are summarized in Tables 2 to 8.

Sampling station No. 1 (Canale Caterattino) showed a high density of bacterial indicators (total and faecal coliforms and faecal streptococci) only in water samples collected in August. The other water samples showed acceptable levels of bacterial indicators. This conforms with the seasonal presence of small discharges (see 2.2). The sand samples showed a constant level of bacterial indicators, which were subject to no fluctuation during the period of study. The DO concentration was at acceptable levels with the exception of the sample collected in August, this correlating with the density of bacterial indicators in the water samples.

Sampling station No. 2 (beach, km 25.6) showed very low levels of bacterial indicators in both water and sand samples. The only exception was the relatively high density of faecal streptococci in the water sample collected on 15.6.82. The DO concentration and pH values were at excellent levels in all the samples examined.

Sampling station No. 4 (Rio Martino) showed very high levels of bacterial indicators in all the samples and correspondingly low values of DO. The pH values were acceptable.

Sampling station No. 5 (beach, km 17.1) showed relatively high levels of bacterial indicators only in the sample collected on 15.6.82. The levels however did not exceed the limits prescribed by the Italian regulations for bathing (100 FC/100 ml). The values of DO and pH were satisfactory in all the samples.

The levels of bacterial density in sand samples were acceptable, though levels found in sampling station No. 2 indicated the influence of the pollution source at station No. 4.

Sampling station No. 6 showed high levels of bacterial indicators in all the water and sand samples (with the exception of the water sample collected on 13.7.82), exceeding the limits of the Italian regulations. The high levels found were due to the influence of the pollution source at station No. 7. The levels of bacterial density in the sand were very high in all the samples. The values of DO and pH were satisfactory in all the samples.

Sampling station No. 7 (Fosso Mascarello) showed very high levels of bacterial indicators in all the samples, which did not correlate to the DO values. The pH values were acceptable.

Sampling station No. 8 showed very high levels of bacterial indicators in the water samples collected on 1.6.82, 13.7.82 and 10.8.82; medium levels in the water samples collected on 18.5.82; and low levels in samples collected on 15.6.82, 26.6.82, 27.7.82 and 31.8.82. Levels of DO and pH were satisfactory. All the sand samples showed a relatively high density of bacterial indicators.

4. Discussion

The levels of the various parameters are shown in Figure 3, as measured in the first sampling site in eight successive weeks. The total coliform (TC), faecal coliform (FC) and faecal streptococci (FS) levels in water samples (the first three graphs in Figure 3) appear to show comparable time trends. The TC, FC and FS levels for sand appear more stable than the corresponding levels in seawater (variances of log-transformed TC and FC values in the sand were significantly less than the corresponding ones in the seawater samples). Analogous results have been recorded in other consistently contaminated sites. It is considered that further data would be necessary to confirm this.

Figure 4 shows contamination levels as measured in five sampling sites three successive weeks. The various indicators generally assume maximum and minimum values in the same sites, thus showing that they basically record the same contamination phenomenon. Significant correlations were found among the various parameters.

Figure 5 shows a highly significant correlation between the FC (MPN) and FC (MF) values. The two parameters appear to be basically equivalent (on the average, MF values were somewhat lower than MPN values, in particular when contamination levels were low). Log-transformed values have been considered, because such transformation enabled a Gaussian-like statistical distribution to be obtained, thus simplifying the statistical treatment of data (Geldreich, 1975).

Figure 6 shows a highly significant correlation between log-transformed FC (MPN) data and log-transformed TC (MPN) data, and Figure 7 the correlation between log-transformed FC (MF) and TC (MPN) data. Figure 8 shows the correlation between log-transformed FS (MF) and FC (MF) log data. Figure 9 shows the correlation between the TC and FC (MPN) log-values in the seawater and in the corresponding sand samples respectively.

All the data obtained indicate that the TC, FC and FS levels in the seawater and in the sand samples are generally correlated, thus indicating that these parameters basically obtain a measure of the same pollution level. The high variability of levels measured in successive samplings is also evident. In any case, the data shown allow an easy classification of site contamination level, if several measurements are considered, and the mean value is assumed as the best indicator (as a rule, the geometric mean or the log-mean or the median value). Single values are scarcely significant for this purpose. The measurement repeatability, as tested by the means of replicated analyses on the same samples, appeared to be characterized by a log-data standard deviation (napierian logarithms) of about 0.7 - 0.8, corresponding to about a factor of 2 for original data in the case of TC and FC, and slightly better for FS. This variability may account only for a minor part of overall variability of measured values.

 $\begin{tabular}{l} \hline \textbf{Table 1} \\ \hline \textbf{Sea and meteorological conditions on the sampling day} \\ \hline \end{tabular}$

Sea conditions	Surface current m/min	Wind	Sky	Last relevant precipitation
calm	10 from E	weak from SW	clear	+ 4 days
calm	absent	weak from E	clear	+ 4 days
rough	absent	strong from E	clear	2 days
slightly rough	absent	weak from E	clear	+ 4 days
calm	3 from E	absent	clear	+ 4 days
slightly rough	absent	weak from E	clear	+ 4 days
calm	1 from E	weak from E	overcast	2 days
calm	20 from W	weak from W	clear	+ 4 days
	calm calm rough slightly rough calm slightly rough calm	calm 10 from E calm absent rough absent slightly rough absent calm 3 from E slightly rough absent calm 1 from E	calm 10 from E weak from SW calm absent weak from E rough absent strong from E slightly rough absent weak from E calm 3 from E absent slightly rough absent weak from E calm 1 from E weak from E	calm 10 from E weak from SW clear calm absent weak from E clear rough absent strong from E clear slightly rough absent weak from E clear calm 3 from E absent clear slightly rough absent weak from E clear calm 1 from E weak from E clear

Table 2

Monitoring results: sampling station No. 1

			Surface w	ater/100	ml	Sa	md/100 g	FELL A	Optio parame		Oth param	er eters
Date	Hour	Total coli. MPN	Faecal coli. MPN	Faecal coli. MF	Faecal strep. MF	Total coli. MPN	Faecal coli. MPN	Faecal strep. MF	DO mg/l	рН	water temp. °C	air temp °C
18.5.82	11.40	1600	350	300	70	230	90	70	10.5	8.3	23	20
1.6.82	11.30	1600	33	40	93	1100	95	10	7.0	7.9	26	29
15.6.82	9.30	70	49	51	39	210	70	20	10.7	8.3	21	22
26.6.82	9.30	23	23	11	58	210	70	0	7.2	8.3	26	22
13.7.82	9.30	14	14	19	16	230	230	40	6.8	8.2	27	28
27.7.82	9.00	46	46	58	10	430	90	40	6.9	8.2	27	29
10.8.82	12.30	>2400	1600	>300	>300	90	40	10	4.3	7.8	26	30
31.8.82	14.20	>2400	>2400	>300	>300	150	90	10	1.0	7.2	28	29

Table 3

Monitoring results: sampling station No. 2

D =	.,		Surface w	ater/100	m1	Sa	nd/100 g		Optio parame		Oth param	er eters
Date	Hour	Total coli. MPN	Faecal coli. MPN	Faecal coli. MF	Faecal strep. MF	Total coli. MPN	Faecal coli. MPN	Faecal strep. MF	DO mg/1	рН	water temp. °C	air temp °C
18.5.82	12.00	5	< 2	0	0	0	0	10	11.5	8.1	23	20
1.6.82	11.40	4 2	< 2	0	86	0	0	0	11.0	8.3	23	29
15.6.82	9.50	11	11	15	>300	40	30	30	13.7	8.4	17	22
26.6.82	9.55	8	8	5	76	0	0	0	11.4	8.2	22	22
13.7.82	9.45	< 2	< 2	0	7	150	90	30	9.8	8.2	27	28
27.7.82	9.15	2	2	0	7	90	40	30	11.3	8.1	22	29
10.8.82	12.50	2	2	0	5	0	0	0	10.0	8.3	27	29
31.8.82	14.40	<2	<2	0	4	0	0	0	9.7	8.3	27	29

Table 4

Monitoring results: sampling station No. 4

Data			Surface	water/100	m1	Sa	ind/100 g		Option parame		Oth param	er eters
Date	Hour	Total coli. MPN			Faecal strep. MF	Total coli. MPN	Faecal coli. MPN	Faecal strep. MF	DO mg/1	рН	water temp. °C	air temp. °C
18.5.82	12.30	23000	13000	8000	13000	====	**==		1.0	7.2	21	20
1.6.82	12.15	70000	70000	51000	1000	*===	====	5445	1.2	7.0	26	24
15.6.82	10.15	1600000	1600000	>300000	6000	====		====	1.0	7.3	22	22
26.6.82	10.20	240000	93000	110000	1000	====	3 33 4	====	0.8	7.6	25	25
13.7.82	10.15	1600000	1600000	>300000	13000	====	====		0.6	7.2	26	29
27.7.82	9.45	1600000	43000	28000	13000	====	====		0.8	7.4	26	28
10.8.82	11.50	>2400000	430000	>300000	45000	====		====	0.9	7.2	27	28
31.8.82	13.55	>2400000	>2400000	>300000	21000		====		0.6	7.5	25	28

			Surface w	ater/100	ml	Sa	nd/100 g		Optio parame		Oth param	er eters
Date	Hour	Total coli. MPN	Faecal coli. MPN	Faecal coli. MF	Faecal strep. MF		Faecal coli. MPN	Faecal strep. MF	DO mg/l	рН	water temp. °C	air temp °C
18.5.82	12.35	240	79	46	3	90	3	80	8.0	8.0	23	22
1.6.82	12.25	43	23	27	40	>11000	4600	40	9.7	8.2	26	26
15.6.82	10.35	150	110	86	>300	150	70	30	11.3.	8.1	18	22
26.6.82	10.35	<2	<2	0	20	90	42	40	9.1	8.3	22	25
13.7.82	10.30	< 2	<2	0	38	90	40	10	7.4	8.2	27	27
	10.00	34	34	39	24	230	230	40	9.5	8.2	22	26
27.7.82	11.30	33	26	6	13	90	40	10	8.3	8.2	26	27
10.8.82 31.8.82	13.40	<2	< 2	5	15	10	10	10	11.0	8.4	27	28

Table 6

Monitoring results: sampling station No. 6

*			Surface w	ater/100	m1	S	and/100 g	, ,	Optio parame		Oth param	er eters
Date	Hour	Total coli. MPN	Faecal coli. MPN	Faecal coli. MF	Faecal strep. MF	A PARK TO	Faecal coli. MPN	Faecal strep. MF	DO mg/l	рН	water temp. °C	air temp °C
18.5.82	13.00	350	70	3	6	>11000	>11000	900	9.0	8.1	24	22
1.6.82	12,40	>2400	>2400	>300	5	640	390	500	10.8	8.2	26	24
15.6.82	10.45	>2400	1600	>300	100	1500	280	80	11.1	8.3	25	22
26.6.82	11.00	1600	1600	210	50	>24000	280	80	8.2	8.1	27	25
13.7.82	11.00	46	46	12	45	>24000	280	250	8.1	8.2	29	27
27.7.82	10,10	350	350	>300	>300	>24000	4600	520	8.3	8.3	28	28
10.8.82	11.00	350	350	180	> 300	4600	200	80	7.9	8.2	28	27
31.8.82	13.15	>2400	>2400	>300	82	1500	390	28	10.6	8.3	28	27
					iğe in	s 1 or 5		-				

Table 7

Monitoring results: sampling station No. 7

Date	Hour		Surface	water/10	00 ml	Sa	and/100 g		Optio parame		Oth param	ner meters
Date	11001	Total coli. MPN	Faeca coli MPN			Total coli. MPN	Faecal coli. MPN	Faecal strep. MF	DO mg/l	рΗ	water temp. °C	air temp °C
18.5.82	13.05	>2400000	1600000	>300000	13000	====	====	2022	10.5	7.8	20	22
1.6.82	12.50	160000	160000	205000	1000	====	=	====	10.1	7.8	21	25
15.6.82	11.00	33000	33000	22000	2000	====	====	====	11.3	7.9	19	22
26.6.82	11.20	350000	240000	210000	1000		====	====	10.4	7.8	21	25
13.7.82	11.30	33000	33000	15000	900	2323	====	====	9.3	7.9	21	28
27.7.82	10.30	2400000	2400000	>300000	1000	====	====	====	9.0	7.6	22	29
10.8.82	10.40	>2400000	350000	>300000	>300000	322	====	====	10.3	7.7	19	26
31.8.82	13.00	23000	5000	18000	850	34 7 =	4445	72nn	11.2	8.0	21	27

Table 8

Monitoring results: sampling station No. 8

Date	Hour	Vour	Surface water/100 ml					and/100 g	Optional parameters		Other parameters		
	nour		Total coli. MPN	Faecal coli. MPN	Faecal coli. MF	Faecal strep. MF	Total coli. MPN	Faecal coli. MPN	Faecal strep. MF	DO mg/l	pН	water temp. °C	air temp °C
18.5.82	13.20		1600	350	215	14 >	11000	>11000	900	9.4	8.1	27	22
1.6.82	13.00		1600	1600	> 300	>300	11000	95	500	8.8	8.2	30	25
15.6.82	11.15		8	8	0	>300	280	30	80	12.1	8.1	19	23
26.6.82	11.30		17	17	0	32	430	230	80	10.7	8.4	22	25
13.7.82	12.00		350	350	>300	>300	150	90	250	8.5	8.3	29	28
27.7.82	11.00		13	13	0	3	430	430	520	11.4	8.3	23	28
10.8.82	10.20		240	240	180	>300	930	110	80	10.5	8.3	24	23
31.8.82	12,40		< 2	<2	0	0	60	30	9	11.7	8.4	22	27

Figures 1 and 2
Selected sampling stations, Province of Latina

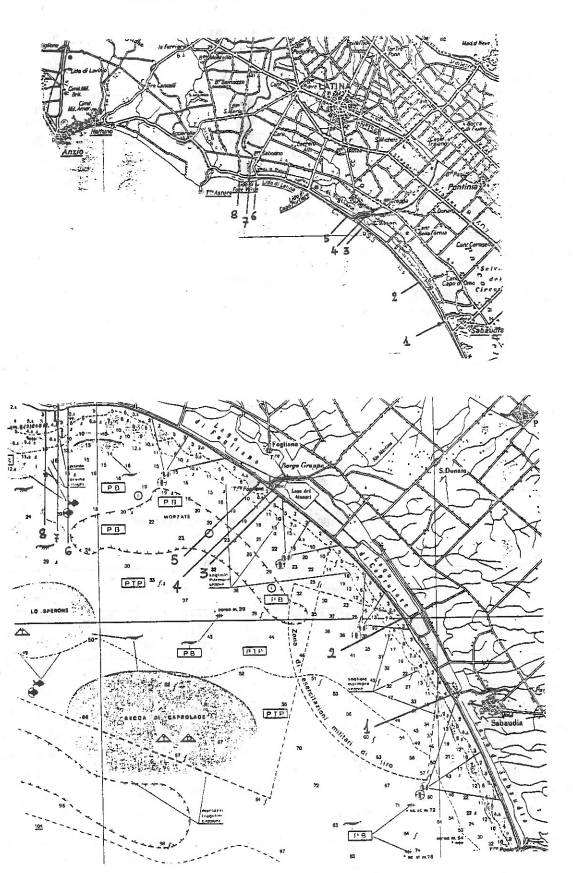
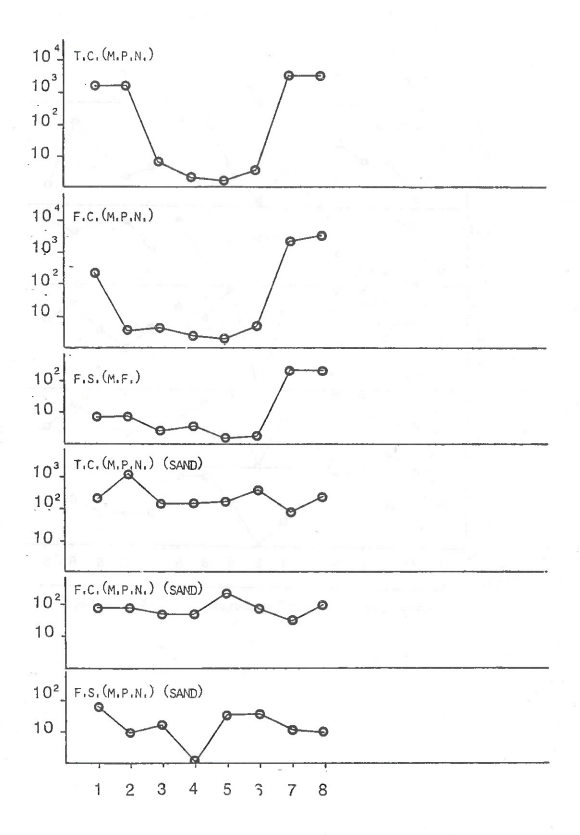
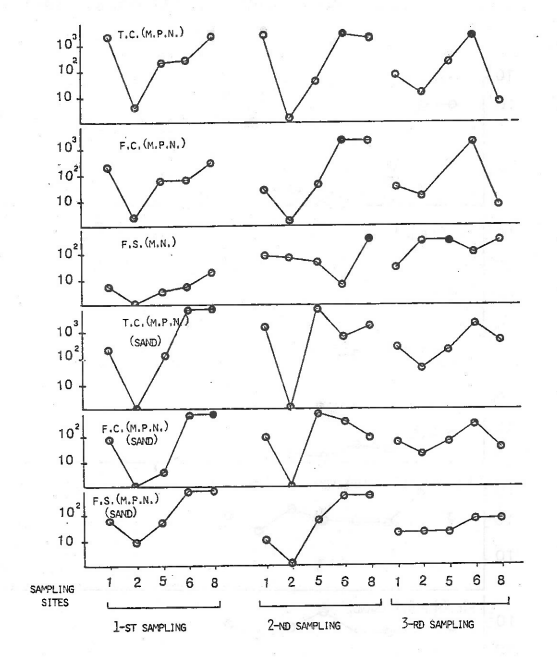


Figure 3

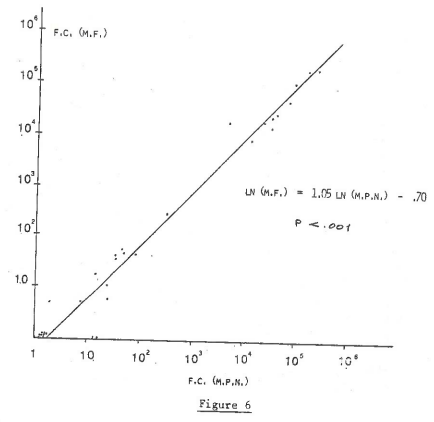
Levels of selected parameters measured at Sampling Site 1 in 8 successive weeks



 $\frac{\text{Figure 4}}{\text{Levels of contamination as measured in 5 sampling sites 3 successive weeks}}$



 $\frac{\text{Figure 5}}{\text{Statistical analysis of FC (MPN) and FC (MF) values}}$



Statistical analysis of log-transformed FC (MPN) data and log-transformed TC (MPN) data

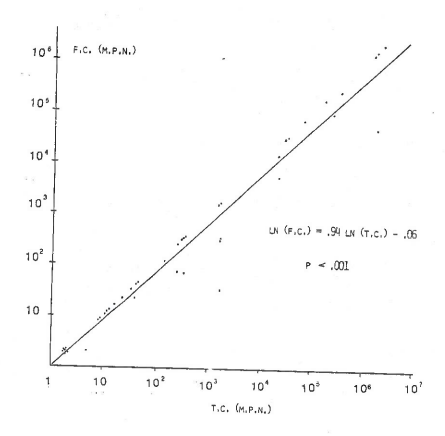


Figure 7

Statistical analysis of log-transformed FC (MF) data and log-transformed TC (MPN) data

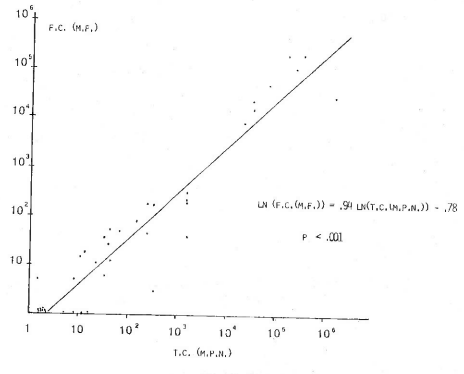
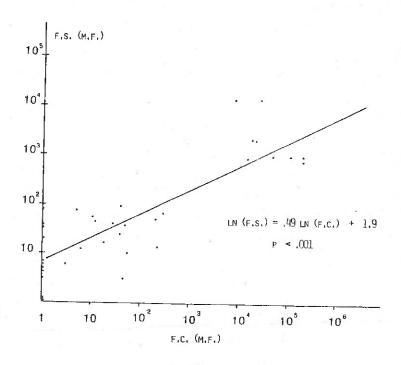
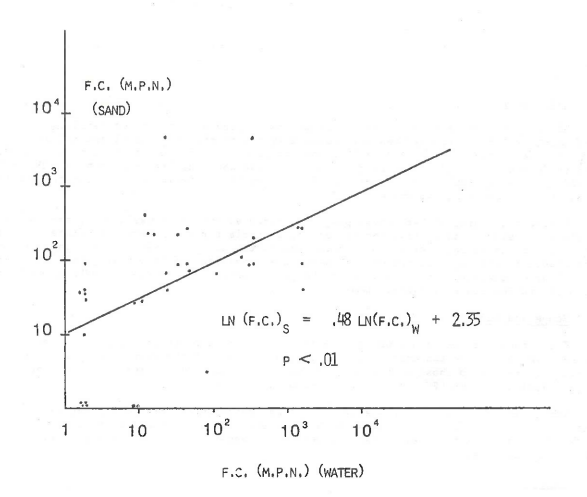


Figure 8

Statistical analysis of log-transformed FS (MF) and log-transformed FC (MF) data



 $\frac{\text{Figure 9}}{\text{Statistical analysis of log-transformed FC (MPN) values in seawater and samples}}$



Annex 6

A COMPARATIVE STUDY OF THE MEMBRANE FILTRATION AND MOST PROBABLE NUMBER METHODS IN THE MICROBIOLOGICAL ANALYSIS OF SEAWATER

by T. Feliu Mendez, S. Grané Terradas and A. Hernández Higuera Regional Government of Catalonia, Tarragona, Spain

1. Introduction

Since 1976, the Territorial Health Promotion Service of the city of Tarragona, Regional Government of Catalonia, has been developing a system of microbiological control of the waters of the province's beaches. In 1978-1979, it participated in the pilot project on Coastal Water Quality Control (MED VII), jointly coordinated by the World Health Organization (WHO) and the United Nations Environment Programme (UNEP), as part of the Mediterranean Pollution Monitoring and Research Programme (MED POL Phase I).

During the course of both these projects, repeated reference was made to the need for unifying sampling procedures, analytical techniques and criteria for interpreting findings. For this reason and on the basis of work performed to date, the Institution expressed its willingness to participate in the study on methods of sampling and analysis of bacteriological parameters in coastal water quality, carried out under the joint sponsorship of WHO and UNEP and coordinated by the Istituto Superiore di Sanità, Rome.

Scope and purpose

The purpose of this study was to compare the membrane filtration and most probable number methods, which are the two most commonly used in analysis of the microbiological quality of coastal waters with regard to those microorganisms indicating faecal contamination, i.e. total coliforms (TC), faecal coliforms (FC) and faecal streptococci (FS).

Three sampling stations were selected. These represented three different levels of faecal contamination, ranging from a coastal area with little or no contamination to an urban wastewater outfall. In each case, repeated samples were taken and analysed microbiologically, using the two methods mentioned above.

Sampling

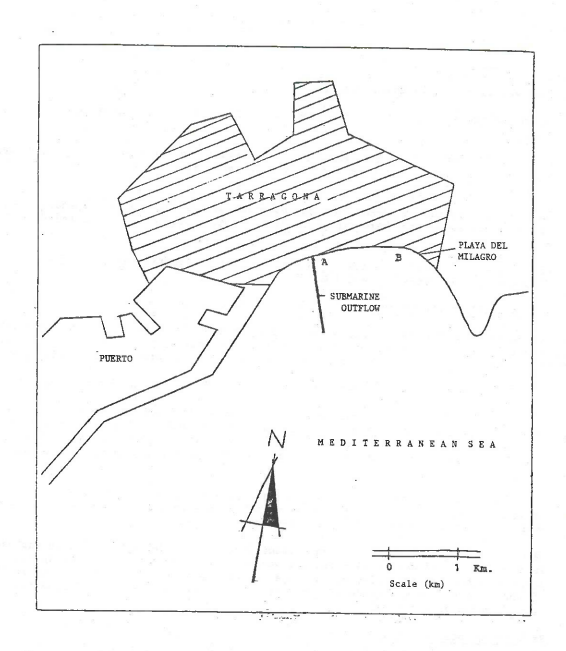
The sampling points selected were:

- (a) the submarine outfall of urban wastewater from the city of Tarragona (length 1000 m, outfall depth 20-25 m);
- (b) the Playa del Milagro, Tarragona, which is affected by submarine outfall A (diagram 1);
- (c) the Playa de la Mora, Tarragona, which is exceptionally clean and hygienic.

Sampling was performed between 10 August and 6 September 1982, most of the samples being taken between 8 a.m. and 10 a.m., using amber glass bottles with ground-glass stoppers, sterilized in a Pasteur oven. At points B and C, sampling was performed by direct access up to 10-15 m from the coastline, at a depth of 1.00-1.50 m, the bottle being dipped 15-20 cm below the surface of the water. Sampling at point A was performed directly at the pumping station for the submarine wastewater outfall.

As soon as samples were collected, they were placed in a thermally insulated container and protected from the light until they reached the laboratory. The time between the collection of the first sample and its arrival at the laboratory was in all instances less than two hours, and the samples were analysed immediately.

Diagram 1. Situation of sampling points A and B (Playa del Milagro, Tarragona)



4. Analytical techniques

The analytical techniques used were those described by APHA (1975), WHO/UNEP (1977) and UNEP (1981).

In addition to the comparison made be ween the membrane filtration and most probable number methods, analysis of TC and FC using the membrane filtration method was carried out in duplicate, using absorbent pads and agar respectively, thus giving a third set of results for these microorganisms.

The use of Escherichia coli (EC) as an indicator microorganism was regarded for the purposes of evaluating the water quality of a coastal area as practically the equivalent of FC.

4.1 Membrane filtration method

For analysis of the three categories of microorganisms of the indicator type, membranes used were Millipore Standard HAWG, 47 mm in diameter and 0.45 µm in pore size. Filtration funnels and supports were made of plastic and sterilized with ultraviolet rays. Culture media were prepared daily, and the volume of water filtered was decided in the light of the concentration of microorganisms anticipated, so as to make the resulting number of colonies 20-80 per plate.

The average number of filtrations per sample of water analysed was eight: three for TC, three for FC, and two for FS. In the case of highly polluted water samples, a series of prior dilutions was carried out, using distilled water, sterile and stoppered.

Culture media and incubation conditions used for each of the indicator microorganisms were as follows.

- (1) Total coliforms: M-Endo broth MF on absorbent pads or M-Endo MF on agar, depending on whether pads or agar were used, on a hermetic petri dish, 47 mm in diameter, incubated for 24 hours at $36 \pm 1\,^{\circ}\text{C}$.
- (2) Faecal coliforms: M-FC broth on absorbent pads or M-FC agar, depending on whether pads or agar were used, on a hermetic petri dish, 47 mm in diameter, incubated for 24 hours at 44 ± 0.2 °C.
- (3) Faecal streptococci: Agar-M-Enterococci, on a hermetic petri dish, 47 mm in diameter, incubated for 48 hours at 36 \pm 1 $^{\circ}$ C.

4.2 Most probable number method

For analysis of the three indicator microorganisms, four series of five tubes were used to determine the most probable number of microorganisms contained in each sample of water. For the analysis of samples of highly polluted water, a sterile phosphate plug was used as a dilutant:

The culture media and the incubation conditions for each of the microorganisms were as follows.

- (1) Total coliforms: MacConkey broth incubated at $36 \pm 1^{\circ}\text{C}$ for 48 hours. Tubes were considered positive when they turned the medium over and produced gas in a Durham tube. Thus, the presumptive number of coliform bacteria was obtained. The positive tubes were redispensed in brilliant green broth at $36 \pm 1^{\circ}\text{C}$ for 48 hours, giving the confirmatory number of coliform bacteria.
- (2) Faecal coliforms: MacConkey broth incubated at $36 \pm 1^{\circ}\text{C}$ for 48 hours. The reading was taken as in determining TC. The positive tubes were redispensed in brilliant green broth at $44 \pm 0.2^{\circ}\text{C}$ for 24 hours, thus giving the number of FC.
- (3) Faecal streptococci: for the presumptive test, azide dextrose broth was used, incubated at $36 \pm 1^{\circ}\text{C}$ for 48 hours. The reading was taken by turbidity. The redispensing of the positive tubes was done with ethyl violet azide broth (EVA broth) incubated at $36 \pm 1^{\circ}\text{C}$ for 48 hours. Tubes were considered positive if there was turbidity of the medium and a purple bead appeared at the base.

Results and statistical analysis

The results obtained from microbiological analyses of TC, FC and FS found at the three sampling points, carried out in accordance with the two methods, can be seen in Tables 1, 2 and 3. The number of samples taken at the submarine outfall was greater because of the variability of the microbiological characteristics of the wastewater. In the case of TC and FC, the results of the analyses made using the membrane filter method, with absorbent pads instead of agar, were also added.

For a statistical comparison of the series of microbiological results obtained from each water sample using the two methods, a parametric statistical method of "comparison of pairs" known as the "t-test" was applied. In order to apply this "t-test", the two series of microbiological results to be compared have to be regarded as adjusting to a normal distribution, which is actually the case if the logarithm of the microbiological concentrations obtained is considered.

Table 1. Results of microbiological analyses carried out at Playa del Milagro, submarine outfall, point A

Date	Temperature	Total coliforms are per 100 ml				eal colifor per 100 ml	Faecal streptococci per 100 ml		
6		MF (pads)	MF (agar)	MPN	MF (pads)	MF (agar)) mpn	MF (agar) MPN
10.8.82 11.8.82 12.8.82 16.8.82 23.8.82 24.8.82 30.8.82 31.8.82 6.9.82	24.8° 25.0° 25.0° 26.5° 25.5° 24.5° 24.8° 24.0° 25.5° 25.4°	3.0.10 ⁸ 9.6.10 ⁷ 1.2.10 ⁸ 1.4.10 ⁷ 4.0.10 ⁷ 2.3.10 ⁷ 5.2.10 ⁷ 1.6.10 ⁷ 3.4.10 ⁷ 2.6.10 ⁷	4.0.10 ⁸ 2.4.10 ⁷ 6.0.10 ⁷ 1.7.10 ⁷ 6.3.10 ⁷ 2.4.10 ⁷ 7.2.10 ⁷ 3.5.10 ⁷ 3.6.10 ⁷ 2.0.10 ⁷	5.0.10 ⁷ 2.0.10 ⁵ 2.4.10 ⁷ 2.4.10 ⁷ 9.2.10 ⁷ 3.4.10 ⁶ 1.6.10 ⁸ 7.9.10 ⁶ 3.5.10 ⁷ 2.4.10 ⁷	5.5.10 ⁷ 6.0.106 ≥ 105 1.1.10 ⁷ 3.4.10 ⁷ ≥ 105 3.4.10 ⁷ 8.4.10 ⁶ 1.7.10 ⁷ 4.5.10 ⁶	8.0.10 ⁷ 1.0.10 ⁵ > 10 ⁵ 1.1.10 ⁷ 2.2.10 ⁷ > 10 ⁵ 1.2.10 ⁷ 1.0.10 ⁶ 1.1.10 ⁷ 8.0.10 ⁶	2.0.10 ⁷ > 10 ⁵ 2.0.10 ⁵ 2.4.10 ⁷ 2.0.10 ⁵ 2.0.10 ⁵ 1.7.10 ⁶ 7.9.10 ⁶ 1.4.10 ⁶ 9.0.10 ⁵	> 10 ⁵ 1.1.10 ⁶ 1.9.10 ⁶ 5.6.10 ⁵ 2.6.10 ⁶ 1.8.10 ⁶ 6.4.10 ⁵ 3.7.10 ⁶ 9.6.10 ⁵ 1.7.10 ⁵	3.3.10 ⁶ 2.0.10 ³ 7.0.10 ⁴ 1.6.10 ⁶ 5.4.10 ⁶ 9.4.10 ⁵ 5.4.10 ⁶ 1.3.10 ⁶ 2.4.10 ⁶ 2.3.10 ⁵

Table 2. Results of microbiological analyses carried out at Playa del Milagro, submarine outfall, point B

Date	Temperature	Total coliforms per 100 ml				coliforms	Faecal streptococci		
		MF (pads)	MF (agar)	MPN	MF (pads)	MF (agar)	MPN	MF (agar)	MPN
10.8.82	24.3	190	130	70	40	5	2	5	2
11.8.82	24.0	100	15	2	5	5	2	31	6
12.8.82	24.6°	700	260	49	12	4	5	7	2
16.8.82	26.0	60	44	26	29	24	21	3	2
24.8.82	24.4	550	46	5	20	7	2	4	/.
30.8.82	24.7°	1100	470	170	110	49	70	8	2
31.8.82	25.0	50	80	9	0	7	4	0	2

Table 3. Results of microbiological analyses carried out at Playa del Milagro, submarine outfall, point C

Date	Temperature		coliforms					Faecal streptococc:		
		MF (pads)	MF (agar)	MPN	MF (pads)	MF (agar)	MPN	MF (agar)	MPN	
10.8.82	24.3°	70	70	2	2	2	2	10		
11.8.82	24.5°	64	44	< 2	0	0	4.0	10	7	
12.8.82	24.4°	94	54	11	,	0	< 2	0	< 2	
16.8.82	26.0°	53	26	7	4.	3	4	3	5	
23.8.82	24.5°	61		,	0	2	< 2	9	6	
24.8.82	24.4		23	2	5	16	< 2	2	< 2	
		280	119	2	2	70	< 2	4	5	
30.8.82	24.3	13	10	< 2	0	6	< 2	0	< 2	
31.8.82	25.0°	44	4	6	2	0	2	0	< 2	

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If the logarithms of the microbiological concentrations obtained using the membrane filter method are represented as:

$$x_1, x_2, \ldots, x_i \ldots, x_n$$

with a mean of $\mu_{\rm X}$ and a standard deviation of $\delta\,x$,

and if the logarithms of the microbiological concentrations obtained using the most probable number method are represented as:

$$y_1, y_2, \ldots, y_i \ldots, y_n$$

with a mean of μ_y and a standard deviation of δ_y^2 ,

then the "t-test" consists in obtaining the difference for each pair of results relating to the same water sample and to the same microorganism (X_i, Y_i) , i.e.:

$$d_{1} = X_{1} - Y_{1}$$

$$d_{2} = X_{2} - Y_{2}$$

$$\vdots$$

$$\vdots$$

$$d_{i} = X_{i} - Y_{i}$$

$$\vdots$$

$$\vdots$$

$$\vdots$$

$$\vdots$$

This series of differences d_1 will also follow a normal distribution whose mean is μ_d , and whose standard deviation is δ_d

Before the two series of results X_i , Y_i can be regarded as not presenting significant differences at level of confidence α , the following hypothesis must hold good:

$$H_0: \mu_d = \mu_X - \mu_Y = 0$$

where

$$\left|\begin{array}{c} u_{\underline{d}} & \sqrt{n} \\ \delta_{\underline{d}} \end{array}\right| < \left|\begin{array}{c} t \\ 1-\alpha, n-1 \end{array}\right|$$

and μ_d is the mean of the differences

δd is the standard deviation

α is the level of confidence

n is the number of differences involved

t is the probability as shown in Table 4.

If H_{0} does not hold good, there are two alternative hypotheses:

(a) where

$$\mu_1 : \mu_d = \mu_X - \mu_Y > 0$$

$$\mu_d \frac{\sqrt{n}}{\delta} > t_{1-\alpha, n-1}$$

i.e. the resulting figures for series X_i (membrane filter method) are higher than those for series Y_i (most probable number method);

(b) where

$$H_1: \mu_d = \mu_X - \mu_Y < 0$$

$$\mu_{\frac{d}{\delta}}^{\frac{d}{\delta}}$$
 < $t_{1-\alpha,n-1}$

i.e. the figures for series Y_i (most probable number method) are higher than those for series X_i (membrane filter method).

Table 4. Probabilities "t" for the application of the "t-test"

The probability of a difference numerically greater than \underline{t} is twice that shown at the head of the table.

Simplified tables of \underline{t} distributions

Degree of freedom			-	Probabil	ity of a	differe	nce grea	ter than	<u>t</u>			
<u>n</u>	.003	.01	.025	.05	.1	.15	.2	.25	.3	.35	.4	.45
1	63.657	31.821	12.706	6.314	3.078	1.963	1.376	1.000	.727	.510	.325	1.5
2	9.925	6.965	4.303	2.920	1.858	1.386	1.061	.816	.617	.445	.289	.15
3	5.841	4.541	3.182	2,353	1.638	1.250	.978	.765	.584	.424	.277	.13
4	4.604	3.747	2.776	2.132	1.533	1.190	.941	.741	.569	.414	.271	.13
5	4.032	3.365	2.571	2.015	1.476	1.156	.920	.727	.559	.408	.267	.13:
6	3.707	3.143	2.447	1.943	1.440	1.134	.906	,718	.553	.404	.265	.13
7	3.499	2.998	2.365	1.895	1.415	1.119	.896	.711	.549	.402	.263	.130
8	3.355	2.896	2.306	1.860	1.397	1.108	.889	.706	.546	.399	.262	.13
9	3.250	2.821	2.262	1.833	1.383	1.100	.883	.703	.543	.398	.261	.12
10	3.169	2.764	2.228	1.812	1.372	1.093	.879	.700	.542	.397	.260	.12
11	3.106	2.718	2.201	1.796	1.363	1.088	.876	.697	.540	.396	.260	.129
12	3.055	2.681	2.179	1.782	1.356	1.083	.873	.695	.539	.395	.259	.128
13	3.012	2.650	2.160	1.771	1.350	1.079	.870	.694	.538	.394	.259	.128
14	2.977	2.624	2.145	1.761	1.345	1.076	.868	.692	.537	.393	.258	.128
15	2.947	2.602	2.131	1.753	1.341	1.074	.866	.691	.536	.393	.258	.128
16	2.921	2.583	2.120	1.746	1.337	1.071	.865	.690	.535	.392	.258	.128
17	2.898	2.567	2.110	1.740	1.333	1.069	.863	.689	.534	.392	.257	.128
18	2.878	2.552	2.101	1.734	1.330	1.067	.862	.688	.534	.392	.257	.127
19	2.861	2.539	2.093	1.729	1.328	1.066	.861	.688	.533	.391	.257	.127
20	2.845	2.528	2.086	1.725	1.325	1.064	.860	.687	.533	.391	.257	.127
21	2.831	2.518	2.080	1.721	1.323	1.063	.859	.686	.532	.391	.257	. 127
22	2.819	2.508	2.074	1.717	1.321	1.061	.858	.686	.532	.390	.256	.127
23	2.807	2.500	2.069	1.714	1.319	1.060	.858	.685	.532	.390	.256	.127
24	2.797	2.492	2.064	1.711	1.318	1.059	.857	.685	.531	.390	.256	.127
25	2.787	2.485	2.060	1.708	1.316	1.058	.856	.684	.531	.390	.256	. 127
26	2.779	2.479	2.056	1.706	1.315	1.058	.856	.684	.531	.390	.256	.127
27	2.771	2-473	2.052	1.703	1.314	1.057	.855	.684	.531	.389	.256	.127
28	2.763	2.467	2.048	1.701	1.313	1.056	.855	.683	.530	.389	.256	.127
29	2.756	2.462	2.045	1.699	1.311	1.055	.854	.683	.530	.389	.256	,127
30	2.750	2.457	2.042	1.697	1.310	1.055	.854	.683	.530	.389	.256	.127
00	2.576	2.326	1.960	1.645	1.282	1.036	.842	.674	.524	.385	.253	.126

Source: Hoel, P.G. A first course in the theory of modern statistical methods. New York, Wiley, 1963 (28).

Table 4 taken from Statistical methods for research workers, with the kind permission of the author, Professor R.A. Fisher, and the publ shers, Oliver and Boyd.

On applying the method to the results obtained from the microbiological analyses with a confidence interval α = 95%, the following results are obtained.

Sampling point A - submarine outfall

(1) Total coliforms: MF (agar)/MPN

$$n = 10$$
 $\mu_d = 0.415$
 $\delta_d = 0.737$
 $t = 2.262$

$$\left| \frac{0.415. \sqrt{10}}{0.737} \right| = 1.779 < 2.262$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(2) Total coliforms: MF (pads)/MPN

$$n = 10$$
 $\mu_d = 0.423$
 $\delta_d = 0.925$
 $t = 2.262$

$$\left| \frac{0.423. \sqrt{10}}{0.925} \right| = 1.446 < 2.262$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(3) Total coliforms: MF (pads)/MF (agar)

$$n = 10$$
 $\mu_d = 0.0087$
 $\delta_d = 0.271$
 $t = 2.262$

$$\left| \frac{0.0087. \sqrt{10}}{0.271} \right| = 0.102 < 2.262$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(4) Faecal coliforms: MF (agar)/MPN

$$n = 7$$
 $\mu_d = 0.586$
 $\delta d = 0.955$
 $\epsilon = 2.447$

$$\left| \frac{0.586. \, \text{V}_7}{0.955} \right| = 1.624 \, < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(5) Faecal coliforms: MF (pads)/MPN

$$\left| \frac{0.777. \sqrt{7}}{0.857} \right| = 2.399 < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(6) Faecal coliforms: MF (pads)/MF (agar)

$$n = 8$$
 $\mu_d = 0.390$
 $\alpha d = 0.672$

$$\left| \frac{0.390. \, \sqrt{8}}{0.672} \right| = 1.641 < 2.365$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(7) Faecal streptococci: MF (agar)/MPN

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

Sampling point B - Playa del Milagro

(1) Total coliforms: MF (agar)/MPN

$$\begin{array}{ccc}
n &=& 7 \\
\mu_{d} &=& 0.636 \\
\alpha_{d} &=& 0.319 \\
t &=& 2.447
\end{array}
\qquad \left| \begin{array}{c}
0.636. \ \sqrt{7} \\
0.319
\end{array} \right| = 5.273 > 2.447$$

Hypothesis H_0 does not hold good, and we apply the terms of hypothesis H_1 :

$$t = 1.943$$
 $\frac{0.636. \sqrt{7}}{0.319} = 5.273 > 1.943$

i.e. the figures for the series representing the membrane filter method are higher than those for the series using the most probable number method.

(2) Total coliforms: MF (pads)/MPN

Hypothesis H_O does not hold good, and we apply the terms of hypothesis $\mathrm{H}_1\colon$

$$t = 1.943$$
 $\frac{1.035. \sqrt{7}}{0.634} = 4.318 > 1.943$

i.e. the figures for the series representing the membrane filter method are higher than those for the series using the most probable number method.

(3) Total coliforms: MF (pads)/MF (agar)

$$\begin{vmatrix}
n = 7 \\
\mu_{d} = 0.399 \\
\delta_{d} = 0.434 \\
t = 2.447
\end{vmatrix} = 2.432 < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of

(4) Faecal coliforms: MF (agar)/MPN

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(5) Faecal coliforms: MF (pads)/MPN

$$\begin{array}{ll}
n = 7 \\
\mu_{d} = 0.402 \\
\delta_{d} = 0.617 \\
t = 2.447
\end{array}$$

$$\left| \frac{0.402. \sqrt{7}}{0.617} \right| = 1.723 < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of

(6) Faecal coliforms: MF (pads)/MF (agar)

$$\begin{array}{ccc}
n &= 7 \\
\mu_d &= 0.203 \\
\delta_d &= 0.548 \\
t &= 2.447
\end{array}
\qquad \left| \begin{array}{c}
0.203 \cdot \sqrt{7} \\
0.548
\end{array} \right| = 0.979 < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(7) Faecal streptococci: MF (agar)/MPN

$$\begin{array}{ccc}
n &= 7 \\
\mu_{d} &= 0.305 \\
\delta d &= 0.364 \\
t &= 2.447
\end{array}
\qquad \left| \begin{array}{c}
0.305 \cdot \sqrt{7} \\
0.364
\end{array} \right| = 2.217 < 2.447$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

Sampling point C - Playa de la Mora

(1) Total coliforms: MF (agar)/MPN

Hypothesis H_0 does not hold good, and we apply the terms of hypothesis H_1 :

t = 1.895
$$\frac{0.938. \sqrt{8}}{0.626} = 4.237 > 1.895$$

i.e. the figures for the series representing the membrane filter method are higher than those for the series using the most probable number method.

(2) Total coliforms: MF (pads)/MPN

$$\begin{array}{ccc}
n = 8 \\
\mu_{d} = 1.271 \\
\delta_{d} = 0.476 \\
t = 2.365
\end{array}$$

$$\frac{1.271. \sqrt{8}}{0.476} = 7.551 > 2.365$$

Hypothesis H_0 does not hold good, and we apply the terms of hypothesis H_1 :

t = 1.895
$$\frac{1.271. \sqrt{8}}{0.476} = 7.551 > 1.895$$

i.e. the figures for the series representing the membrane filter method are higher than those for the series using the most probable number method.

(3) Total coliforms: MF (pads)/MF (agar)

$$\begin{vmatrix}
n = 8 \\
u_d = 0.333 \\
\delta_d = 0.318 \\
\epsilon = 2.365
\end{vmatrix} = 2.961 > 2.365$$

Hypothesis Ho does not hold good, and we apply the terms of hypothesis H1:

$$t = 1.895$$

$$\left| \frac{0.333. \, \forall 8}{0.318} \right| = 2.961 > 1.895$$

i.e. the figures for the series representing the membrane filter method using pads are higher than those for the same method using agar.

(4) Faecal coliforms: MF (agar)/MPN

$$\begin{vmatrix}
n = 8 \\
u_d = 0.275 \\
\delta_d = 0.657 \\
t = 2.365
\end{vmatrix} = 0.275 \cdot \sqrt{8} = 1.184 < 2.365$$

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(5) Faecal coliforms: MF (pads)/MPN

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(6) Faecal coliforms: MF (pads)/MF (agar)

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

(7) Faecal streptococci: MF (agar)/MPN

i.e. the two series of results do not present significant differences with a confidence interval of 95%.

The entire statistical analysis of the results is summarized in Tables 5 and 6.

Table 5. Summary comparison of the results of microbiological analysis obtained using the MF and MPN methods

Sampling	Faecal	Total coliforms	Faecal coliforms	Faecal streptococci
point A	contamination High	10 ⁸ - 10 ⁷ MF = MPN	10 ⁷ - 10 ⁶ MF = MPN	$10^6 - 10^5$ MF = MPN
В	Lower	10 ³ - 10 ² MF > MPN	$10^2 - 10$ $MF = MPN$	10 - 0 MF = MPN
С	Minimal	$10^2 - 10$ MF > MPN	10 - 0 MF = MPN	10 - 0 $MF = MPN$

N.B. The concentrations of microorganisms are expressed in 100 ml of the sample.

Thus, the series of results of the microbiological analyses obtained by the MF and MPN methods do not present significant differences with a confidence interval of 95%, except in the case of lower concentrations of TC, where the figures obtained by the MF method are slightly higher than those using the MPN method.

Table 6. Summary comparison of results obtained from analyses of total coliforms and faecal coliforms by the MF method, using absorbent pads or agar

Sampling point	Faecal contamination	Faecal coliforms	Faecal streptococci
A	High	$10^8 - 10^7$ Pads = agar	$10^7 - 10^6$ Pads = agar
В	Lower	$10^3 - 10^2$ Pads = agar	$10^2 - 10$ Pads = agar
С	Minimal	$10^2 - 10$ Pads > agar	10 - 0 Pads = agar

N.B. The concentrations of microorganisms are expressed in 100 ml of the sample.

Thus, the series of results of analyses of TC and FC obtained by the MF method using pads or agar do not present significant differences with a confidence interval of 95%, except in the case of lower concentrations of TC, where the use of pads gives slightly better results than those obtained when agar is used.

6. Comparative economic assessment

With a view to making a comparative economic assessment of the microbiological analyses of the three indicator microorganisms, according as one or other method is used, the following factors were taken into account:

- (1) cost of culture media, reagents and dilutants;
- (2) cost of disposable material;
- (3) cost of depreciation on the inventory;
- (4) time spent in preparing and carrying out the analyses.

This last factor was not included in the cost estimates. A breakdown of costs is given in Tables 7-12.

A summary of the economic assessment is given in Table 13.

Table 7. Average cost of culture media per tube or plate

Culture medium	f	ce per lask setas)	Number o plates p	f tubes/ er flask	tube	cost per e/plate esetas)
			Double	Single	Doub1e	Single
EVA broth	7	539	648	1 297	11.6	5.8
Azide dextroze broth	7	000	648	1 297	10.8	5.4
MacConkey broth	7	300	648	1 297	11.26	5.6
Brilliant green	5	131	648	1 297	7.92	3.96
M-FC broth Agar (FC)	7	670	6 1	91		1.2
M-Endo broth	8	031	1 1	79	· (6	6.8
Agar (TC)		-	***			8.1
M-Enterococcus agar Bacto agar		175 884	1 5 3 6			6.7 1.3

Table 8. Average cost of culture per sample analysed

# g		Total coliforms	Faecal coliforms	Faecal streptococci	Total cost (pesetas)
MF	Number of plates	3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	3	2	1 - 1
-	Average cost	24.3	7.5	13.4	45.2
	Number of tubes (presumptive)	20	<u>-</u>	20	* 1_*
MPN	Number of tubes (confirmed)	20	20	20	
	Average cost	252	79.2	251	582.2

Table 9. Average cost of disposable material per sample of water analysed

Total		524.8 pesetas
	36.0 pesetas per membrane 8 membranes per sample	288 pesetas per sample
Membranes	Use and discard	
	29.6 pesetas per plate 8 plates per sample	236.8 pesetas per sample
Plates	Use and discard	
MF method		
Total		16.8 pesetas
	25 pesetas per D. tube 60 D. tubes per sample	3.8 pesetas per sample
Durham tubes	Average lifetime 1.5 years	
	51 pesetas per tube 100 tubes per sample	13 pesetas per sample
Tubes	Average lifetime 1.5 years	
MPN method		**

Table 10. Cost of depreciation of apparatus (MPN method)

MPN method				
Incubator	(stove)	37°C	Cost: Average lifetime (depreciation): Average number of analyses per week: Average depreciation cost:	43 227 pesetas 10 years 7 33.2 pesetas
Incubator	(stove)	44°C	Cost: Average lifetime (depreciation): Average number of analyses per week: Average depreciation cost:	43 227 pesetas 10 years 7 16.6 pesetas
Autoclave			Cost: Average lifetime (depreciation): Number of times used per analysis: Average number of analyses each week: Average depreciation cost:	400 000 pesetas 10 years 2 7 153.0 pesetas
Total dep	reciatio	n cost	(average)	202.8 pesetas

Table 11. Cost of depreciation of apparatus (MF method)

Total dans	eciation cost (
, 1, 1		Average lifetime (depreciation): Average number of analyses per week: Number of times used per analysis: Average depreciation cost:	400 000 pesetas 10 years 56 1 76.5 pesetas
Autoclave	(stove) 44 C	Cost: Average lifetime (depreciation): Average number of analyses per week: Average depreciation cost: Cost:	43 227 pesetas 10 years 56 2.07 pesetas
	(stove) 37°C	Cost: Average lifetime (depreciation): Average number of analyses per week: Average depreciation cost:	43 227 pesetas 10 years 56 2.07 pesetas
	equipment	Cost: Average lifetime (depreciation): Average number of analyses per week: Average depreciation cost:	153 000 peseta 10 years 56 58.8 pesetas

Table 12. Time spent in preparing and carrying out analyses

	*	MF	MPN
Auxiliary staff	Preparation of culture media and cleaning of tubes	2 hours	4 hours
Technical staff	Dispensing Redispensing	1/2 hour	1/2 hours
Total time	Auxiliary staff Technical staff	2 hours 1/2 hour	2 hours

Table 13. Total cost and time spent for carrying out a microbiological analysis

FM	MPN
45.2 pesetas 524.8 pesetas 88.0 pesetas	582.2 pesetas 16.8 pesetas 202.8 pesetas
658.0 pesetas	801.8 pesetas
2 hours 1/2 hour	4 hours 1 hour
2 1/2 hours	5 hours
	45.2 pesetas 524.8 pesetas 88.0 pesetas 658.0 pesetas 2 hours 1/2 hour

Thus, the average cost of a microbiological analysis of TC, FC and FS is higher using the MPN method than using MF.

The time spent in preparing and carrying out microbiological analyses of TC, FC and FS using the MPN method is approximately double that using the MF method.

Conclusions

The following can be concluded from the results obtained.

- (1) The mean concentrations of microorganisms indicating faecal contamination of urban wastewater in the case of a Spanish city with a population of approximately 100~000 are TC (10^8-10^7), FC (10^7-10^6) and FS (10^6-10^5). These concentrations are expressed in 100~ml of the sample.
- (2) The mean concentrations of microorganisms indicating faecal contamination of the waters in the case of a beach affected by a submarine outfall of specific dimensions and characteristics are TC (10^3-10^2) , FC (10^2-10) and FS (10-0). These concentrations are expressed in 100 ml of the sample.
- (3) The mean concentrations of microorganisms indicating faecal contamination of the waters in the case of a virgin beach, in the sense of one not affected by wastewater effluents, are TC (10^2-10) , FC (10-0) and FS (10-0). These concentrations are expressed in 100 ml of the sample.
- (4) The analysis of TC gives values which present no significant differences whether the method is MF or MPN until a certain minimum bacterial concentration has been reached which can be set at 10^3 . For lower concentrations, the MF method gives slightly better results than MPN.
- (5) For the analysis of FC, it is immaterial, as far as obtaining results is concerned, whether the MF or MPN method is used, since the results of the series obtained by either method present no significant difference.
- (6) For the analysis of FS, it is immaterial, as far as obtaining results is concerned, whether the MF or MPN method is used, since the results of the series obtained by either method present no significant difference.
- (7) For the analysis of TC using the MF method, it is immaterial whether a solid culture medium (agar) or a liquid medium (absorbent pads) is used, except after a certain minimum bacterial concentration which may be set at 10^2 . For lower concentrations, the use of a liquid medium (absorbent pads) gives results which are slightly better than those using a solid medium (agar).

- (8) For the analysis of FC using the MF method, it is immaterial whether a solid culture medium (agar) or a liquid (pad) is used, since the results obtained from the series produced by either method present no significant difference.
- (9) The total incubation time for the various microorganisms indicating faecal contamination is less with the MF method than with MPN, which means that the results are obtained more quickly.
- (10) The MPN method requires far more laborious and lengthy preparation of the culture media, in terms of time and materials, than the MF method.
- (11) The time required for carrying out an analysis using the MPN method is twice that needed for the MF method.
- (12) The MF method has a greater degree of precision, since it provides a direct colony count instead of the statistical approximation required for obtaining results by the MPN method.
- (13) The MF method isolates the bacteria from the liquid in which they are suspended, thus making it possible to analyse water samples containing enzymes and other substances which inhibit growth.
- (14) The MF method is definitely indicated for the analysis of highly mineralized waters, which can produce false reactions in MPN liquid media.
- (15) The MF method is not to be recommended for waters with a high suspended matter content, since the matter in suspension retained in the membrane filter inhibits the perfect diffusion of the nutritive substance through its pores to the bacteria deposited on its upper surface and thus has an unfavourable effect on the growth of the colonies.
- (16) The MF method is not to be recommended for waters with a high suspended matter content, since the filters rapidly become choked up and filtering of representative volumes is
- (17) The MF method enables a larger volume of samples to be analysed than MPN.
- (18) The MF method allows for the filtration of samples $\underline{\text{in situ}}$ and the transfer of the membranes to the laboratory in a preserving medium.
- (19) The average total cost of microbiological analysis of TC, FC and FS is higher with the MPN method than with the MF method.

8. Acknowledgements

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Annexe 7

MISE AU POINT DES TECHNIQUES D'ECHANTILLONNAGE ET D'ANALYSE POUR LA SURVEILLANCE DE LA POLLUTION

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Introduction

Conformément au programme de travail concernant la mise au point des techniques d'échantillonnage et d'analyse pour la surveillance de la pollution proposé par l'Institut Supérieur de la Santé, Rome, l'Institut Pasteur de Tunis a procédé, entre le 15 mai et le 15 août 1982, à des prélèvements hebdomadaires d'eau de mer et de sédiments en trois points sur le littoral de la région de Tunis. Les trois points sont les suivants :

- plage d'Hammam-Lif : milieu marin pollué;
- plage de la Marsa : milieu marin contaminé (sous l'influence du milieu pollué à Hammam-Lif);
- plage Raoued : milieu marin propre.

A partir de ces points nous avons prélevé au total 72 échantillons (répartis en trois groupes de 24 échantillons (12 échantillons "eau de mer" et 12 échantillons "sédiments"). Afin d'attribuer statistiquement plus de crédibilité à nos investigations, nous avons doublé le nombre des échantillons, effectuant des prélèvements chaque semaine, au lieu de tous les 15 jours. L'horaire des prélèvements est donné au tableau l.

Les analyses de ces échantillons ont porté sur la détermination du nombre le plus probable/100 ml (NPP) des coliformes totaux, des coliformes fécaux et d'Escherichia coli. Concernant les dénominations NPP des coliformes fécaux et NPP des E. coli, nous admettons comme E. coli les coliformes qui manifestent à la fois la faculté de fermenter, avec dégagement de gaz, le lactose à 44 °C et de produire l'indole à 44 °C, à part leur conformité aux tests JMVIC.

Nous considérons comme "coliformes fécaux" l'ensemble des E. coli et des coliformes positifs au lactose à 44 °C et négatifs à l'indole.

La technique des tubes multiples dont la phase confirmative exige l'épreuve des deux caractères thermophiles (fermentation du lactose avec dégagement de gaz et production d'indole à 44 °C) nous donne la possibilité de différencier le nombre le plus probable des E. coli.

La technique de la membrane filtrante, qui est conçue à la base exclusivement d'un troisième caractère thermophile (notamment la faculté de certaines souches de coliformes qui circulent dans le milieu marin à croître et à proliférer en colonies sur les membranes filtrantes incubées à 44°C) nous donne la possibilité d'établir seulement le nombre le plus probable des coliformes fécaux et ce dans un sens beaucoup plus large, étant donné que la faculté de croissance à 44 °C est plus commune aux coliformes que celle de fermenter le lactose ou de produire l'indole à cette même température.

Si l'objectif est d'établir laquelle des deux techniques est prioritaire en colimétrie du milieu marin, dès maintenant il faut préciser le paramètre (NPP des coliformes totaux, fécaux ou E. coli) qui servira à faire le choix. Il est important aussi de préconiser le procédé standard à appliquer pour la détermination de ce paramètre.

Dans cet esprit, il est utile de mentionner qu'en ce qui concerne la technique des tubes multiples, et notamment le bouillon lactosé utilisé pour effectuer la phase présomptive de cette technique, l'Institut Supérieur de Santé de Rome a recommandé d'appliquer l'un des deux milieux, soit Mac Conkey Broth, soit Lactose Broth, laissant le choix aux participants de l'étude.

Lors du Programme MED POL VII, nous avons préféré utiliser la technique des tubes multiples comme étant, selon notre expérience, plus valable que celle de la membrane filtrante en appliquant simultanément deux bouillons lactosés - le Brilliant Green Lactose Bile Broth (agréé en France) et le Lactose Broth (agréé aux USA).

Nous avons constaté qu'en utilisant le Lactose Broth, les nombres les plus probables des coliformes totaux et des coliformes fécaux étaient beaucoup plus élevés que ceux obtenus par le Brilliant Green Lactose Bile Broth, ce qui exprime que la sensibilité de la technique des tubes multiples est dépendante du type de bouillon utilisé. Afin de confirmer cette constatation et d'arriver à une meilleure compréhension sur la valeur réelle des deux techniques (tubes multiples et membrane filtrante), nous avons effectué dans le cadre de ce projet la colimétrie de chaque échantillon en appliquant la technique des tubes multiples simultanément en trois variantes, en ce qui concerne le type de bouillon lactosé utilisé.

Pour la première variante, nous avons utilisé Mac Conkey Broth (production déshydratée DIFCO), pour la deuxième variante Lactose Broth (production déshydratée DIFCO), pour la troisième variante Brilliant Green Lactose Bile Broth (production déshydratée PASTEUR). Les volumes d'échantillons ensemencés dans ces différents bouillons lactosés sont : 5 tubes de 10 ml, 5 tubes de 1 ml et 5 tubes de 0,1 ml.

Dans le cas de ces trois variantes, la phase "confirmative" a été effectuée par réensemencements dans le bouillon lactosé bilié au vert brillant - épreuve de la fermentation du lactose à 44 °C - et dans l'eau peptonée - épreuve de la production d'indole à 44 °C - à partir de toutes les cultures propres à la phase présomptive qui ont manifesté la fermentation du lactose avec dégagement de gaz à 37 °C.

En ce qui concerne la technique de la membrane filtrante, nous avons utilisé les membranes et l'équipement MILLIPORE, et la gélose Endo (production déshydratée MERIEUX dans le cas des coliformes totaux et la gélose CF DIFCO pour les NPP des coliformes fécaux). Les résultats obtenus sont présentés au tableau 2.

2. Résultats et discussion

Les NPP/100 ml obtenus par la technique des tubes multiples en utilisant simultanément les trois bouillons lactosés, répartis selon la provenance des échantillons, sont présentés dans les tableaux 3, 4 et 5. Selon ces tableaux, 27,77% des échantillons se sont avérés exempts de coliformes en appliquant ces trois variantes.

Il faut noter que dans la majorité des échantillons "positifs", la concentration des coliformes est minime comparée aux normes propres à l'eau de mer polluée. Cela est dû au fait que le déversement de l'eau d'égout, non épurée dans le point de prélèvement choisi comme "pollué", était éliminé immédiatement après le début de cette étude, en conséquence de la mise en exploitation de la station d'épuration des eaux d'égout.

Cette contamination quoique limitée s'est avérée suffisante à l'étude de la sensibilité de la technique des tubes multiples en fonction du type de bouillon lactosé utilisé. Pour démontrer cette dépendance nous avons gradué en trois niveaux (sensibilité élevée, sensibilié moyenne et sensibilité limitée) les trois bouillons lactosés ci-dessus mentionnés. Dans ce but, nous avons comparé les valeurs des NPP des <u>E. coli</u>, des coliformes fécaux et des coliformes totaux propres à chaque échantillon qui a révélé la présence de coliformes.

Le niveau "sensibilité élevée" est attribué au bouillon lactosé qui a manifesté le NPP (exprimé en chiffres) le plus élevé. Sur ce principe sont évalués les niveaux "sensibilité moyenne" et "sensibilité limitée". Les résultats de cette classification sont présentés dans le tableau 6. Compte tenu de ces résultats comparatifs on peut justifier la conclusion que l'utilisation des bouillons lactosés Lactose Broth s'avère la plus favorable à la sensibilité de la technique des tubes multiples en ce qui concerne la mise en évidence de la présence de coliformes et l'évaluation de leur concentration dans le milieu marin (soit l'eau de mer, soit les sédiments).

La priorité du Lactose Broth par rapport au Mac Conkey et Brilliant Green Lactose Bile Broth est bien confirmée surtout par les chiffres propres aux NPP/100 ml des E. coli présentés dans le tableau 7. Cette priorité du bouillon Lactose Broth provient du fait que les E. coli, après avoir quitté leur habitat naturel (l'organisme humain et animal) pour passer dans les effluents, et enfin dans l'eau de mer, ne trouvent pas toujours dans l'environnement des conditions propices à leur survie. Souvent au moment des prélèvements d'eau de mer et de leur ensemencement au laboratoire, ils sont déjà d'une vitalité fort affaiblie. Aussi est-il important d'éviter l'utilisation, à l'étape présomptive de la technique des tubes multiples, des bouillons lactosés sélectifs (Mac Conkey Broth et Brilliant Green Lactose Bile Broth), qui risquent de gêner plus encore la croissance et la prolifération de ces germes et eur mise en évidence au laboratoire. Il faut rappeler que le Lactose Broth offre aux coliformes des conditions de réanimation pendant la phase de latence, à l'image de celles qu'offre l'eau peptonée aux denrées alimentaires pour leur préenrichissement.

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Afin de faire le choix de laquelle des deux techniques (tubes multiples ou membrane filtrante) doit être préparée dans la pratique de la colimétrie du milieu marin, nous avons comparé les données de notre étude portant sur les NPP/100 ml des E. coli à celles concernant la technique de la membrane filtrante. Les données utilisées à cette comparaison sont présentées aux tableaux 8 et 9.

En comparant les valeurs des NPP des coliformes fécaux obtenues par la technique de la membrane filtrante aux valeurs concernant les NPP des $\frac{E.\ coli}{par}$ déterminées par la technique des tubes multiples (il est difficile de les différencier par la technique de la membrane filtrante pratiquée à 44 °C), on constate que par cette technique dans toutes ses variantes à niveaux différents on arrive à déceler la présence des coliformes dans un nombre plus élevé d'échantillons (en pourcentage) que par la technique de la membrane filtrante (voir tableaux 7 et 8).

3. Conclusion

En conclusion des résultats de cette étude, on peut dire que la technique des tubes multiples s'avère plus efficace que celle utilisant la membrane filtrante quant à la détermination du degré de pollution de l'eau, et ce, utilisant l'un quelconque des trois bouillons lactosés. Par ailleurs, il est à remarquer que le Lactose Broth s'est manifesté bien plus sensible que les autres bouillons lactosés lors de la détermination du degré de pollution de l'eau pour les différents coliformes.

Cependant faut-il rappeler que le nombre d'échantillons utilisés à cette étude est trop réduit pour pouvoir s'assurer une conclusion statistique crédible sinon valable sur la supériorité de la sensibilité de la technique des tubes multiples sur celle des membranes filtrantes.

<u>Tableau 1</u>
Horaíre des prélèvements

Date	Р	ollué		ie prélèvement ontaminé		Propre		
de prélèvement	Eau N d			Eau Sédiment N° d'échantillon		Sédiment 'échantillon		
24.05.82	1	2	3	4	5	6		
31.05.82	7	8	9	10	11	12		
10.06.82	13	14	15	16	1.7	18		
17.06.82	19	20	21	22	23	24		
21.06.82	25	26	27	28	29	30		
26.06.82	31	32	33	34	35	36		
07.07.82	37	38	39	40	41	42		
15.07.82	43	44	45	46	47	48		
19.07.82	49	50	51	52	53	54		
27.07.82	55	56	57	58	59	60		
03.08.82	61	62	63	64	65	66		
10.08.82	67	68	69	70	71	72		

Tableau 2

Résultats obtenus par la méthode utilisant les membranes filtrantes (exprimés en nombre de colonies métalliques/100 ml à 44 °C

5 1 11 - 17 0 23 2 29 0 35 0		Contar	miné	Pollué					
	Nombre des colonies à 44 °C	Numéro des échantillons	Nombre des colonies à 44	°c	Numéro des échantillons	Nombre colonies	des à 44	°(
 5	1	3	1		1	4			
	. <u> </u>	9	3		7	3			
	0	15	0		13	1			
		21	-		19	7			
		27	7		25	35			
	0	33	2		31	10			
41	6	39	22		37	-			
47	0	45	26		43	52			
53	0	51	0		49	40			
59	2	57	12		55	40			
65	0	63	12		61	-			
71	5	69	_		67	20			
6	0	4	0		2	0			
12	2	10	- 10		8	20			
18	1	16	18		14	150			
24	1	- 22	56		20	5			
30	0	28	40		26	-			
36	0	34	8		32	-			
42	20	40	-		38	~			
48	4	46	0		44	-			
54	2	52	20		50	ж —			
60	4	58	-		56	и с			
66		64	e -		62	260			
72	_ = =	70	-		68	0			

Tableau 3

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>pollué</u> Nature de l'échantillon : eau de surface

Туре	Formule	des tubes po	sitifs	Nombre le plus probable, 100			
de bouillon utilisé	Coliformes totaux	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli	
Mac Conkey Broth	5-5-2	5-5-2	5-3-0	542	542	79	
Lactose Broth	5-5-3	5-5-2	5-3-1	920	542	109	
Brilliant G.L.B. Broth	5-1-0	0-0-0	0-0-0	33	< 2	< 2	
Mac Conkey Broth	5-5-5	5-3-0	5-3-0	>2400	79	79	
Lactose Broth	5-5-5	5-3-1	5-3-0	>2400	109	79	
Brilliant G.L.B. Broth	5-2-0	4-2-0	3-2-0	49	22	14	
Mac Conkey Broth	5-5-1	5-2-0	5-1-0	348	49	33	
Lactose Broth	5-5-2	5-2-2	5-1-1	548	94	46	
Brilliant G.L.B. Broth	5-1-0	5-1-0	5-1-0	33	33	33	
Mac Conkey Broth	5-5-5	5-5-5	5-5-5	>2400	> 2400	> 2400	
Lactose Broth	5-5-5	5-5-5	5-5-5	>2400	>2400	> 2400	
Brilliant G.L.B. Broth	5-5-5	5-5-5	5-5-4	>2400	> 2400	1600	
Mac Conkey Broth	5-5-2	5-4-2	2-2-0	542	221	9	
Lactose Broth	5-5-2	5-5-2	5-1-1	542	542	46	
Brilliant G.L.B. Broth	5-4-2	5-4-1	3-2-0	221	172	14	
Mac Conkey Broth	5-3-2	5-2-0	5-1-0	109	49	33	
Lactose Broth	5-5-1	4-2-1	4-1-0	348	26	17	
Brilliant G.L.B. Broth	5-3-1	5-1-1	4-1-0	109	46	17	
Mac Conkey Broth	5-5-5	5-5-5	5-5-5	> 2400	>2400	>2400	
Lactose Broth	5-5-5	5-5-2	5-3-2	>2400	542	141	
Brilliant G.L.B. Broth	5-5-5	5-5-5	5-4-5	>2400	>2400	426	
Mac Conkey Broth	5-3-3	0-0-0	0-0-0	175	<2	2	
Lactose Broth	5-4-3	5-3-2	5-1 - 1	278	141	63	
Brilliant G.L.B. Broth	5-1-1	4-1-0	1-1-0	46	17	4	
Mac Conkey Broth	5-5-5	5-5-5	5-5-5	> 2400	> 2400	>2400	
actose Broth	5-5-5	5-5-5	5-5-5	>2400	> 2400	>2400	
Brilliant G.L.B. Broth	5-5-5	5-5-5	4-5-5	>2400	>2400	1600	
Mac Conkey Broth	5-4-4	5-4-3	0-0-0	345	278	< 2	
Lactose Broth	5-5-3	5-5-3	2-1-0	920	920	7	
Brilliant G.L.B. Broth	5-4-4	5-5 - 0	1-5-0	345	240	24	
Mac Conkey Broth	5-4-5	5-4-2	5-3-1	426	221	109	
actose Broth	5-5-5	5-5-5	5-4-1	>2400	>2400	172	
Brilliant G.L.B. Broth	5-4-2	5-4-2	4-2-1	221	221	26	
lac Conkey Broth	5-4-5	5-3-4	4-3-1	426	212	33	
actose Broth	5-3-3	5-2-1	3-2-1	175	70	14	
Brilliant G.L.B. Broth	5-4-1	5-3-1	5-2-0	172	109	49	

Tableau 3 (suite)

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>pollué</u> Nature de l'échantillon : sédiment

Cara all	Formule	des tubes po	sitifs	Nombre le	plus probabl	e, 100	
Type de bouillon utilisé	Coliformes	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli	
		7.00			6.9	-	
ac Conkey Broth	-		5-3-2	542	542	141	
actose Broth	5-5-2	5-5-2	0-0-0	240	7	< 2	
rilliant G.L.B. Broth	5-5-0	2-1-0	0-0-0		7.000	21	
	5-5-4	4-5-4	3-3-3	1600	1600		
lac Conkey broth	5-5-5	5-5-3	5-5-0	>2400	920	240	
actose Broth	5-5-4	5-5-3	5-3-2	1600	920	141	
Brilliant G.L.B. Broth			4-2-0	1600	1600	22	
lac Conkey broth	5-5-4	5-5-4		1600	240	130	
Lactose Broth	5-5-4	5-5-0	5-4-0	542	4	<2	
Brilliant G.L.B. Broth	5-5-2	0-1-1	0-0-0	742		179	
	5-1-1	2-0-0	1-0-0	46	, 5	2	
Mac Conkey broth	5-1-0	5-0-0	3-0-0	33	23	8	
Lactose Broth	5-1-0 5-0-2	3-0-2	3-0-2	43	13	13	
Brilliant G.L.B. Broth	3 - 0-2			<2	₹2	<2	
Mac Conkey broth	0-0-0	0-0-0	0-0-0	• -	<2	< 2	
Lactose Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	\Z		
	2.0.0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Lactose Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0			10	<2	1100
W. O. Jean broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey broth	0-0-0	0-0-0	0-0-0	< 2	<2		
Lactose Broth		0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth		2 2 2	0-0-0	₹2	<2	42	
Mac Conkey broth	0-0-0	0-0-0		8	6	14	
Lactose Broth	1-2-1	1-1-1	1-1-0	<2	<2	< 2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0		0 2 1 14 1	9 10 10 10	- 1
	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey broth	3-0-0	3-0-0	2-0-0	8	8	5	
Lactose Broth	• • /	0-0-0	0-0-0	<2	< 2	< 2	
Brilliant G.L.B. Broth	0-0-0	0-0-0			<2	42	
Mac Conkey broth	0-0-0	0-0-0	0-0-0	<2	<2	42	
Lactose Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth		0-0-0	0-0-0	<2	< 2		_
2 1/2 1/2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		5-5-5	5-5-5	>2400	>2400	>2400	
Mac Conkey broth	5-5-5		3-4-4	>2400	1600	24	
Lactose Broth	5-5-5	4-5-5	0-3-0	253	212	6	
Brilliant G.L.B. Broth	5-3-5	5-3-4	0-3-0			10	
	5-3-2	5-2-2	2-2-1	141	94	12	
Mac Conkey broth	5-2-2	5-1-1	4-1-1	94	46	21	
Lactose Broth	77	4-1-0	1-0-0	17	1 7	2	
Brilliant G.L.B. Broth	n 4-1-0	7 1 0					

Tableau 4

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>contaminé</u> Nature de l'échantillon : eau de surface

Type	Formule	des tubes po	sitifs	Nombre le	plus probabl	e, 100
de bouillon utilisé	Coliformes totaux	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli
Mac Conkey broth	5-5-3	1-0-0	0-0-0	920	2	<2
Lactose Broth	5-5-3	3-1-1	2-0-0	920	14	5
Brilliant G.L.B. Broth	3-0-0	3-0-0	2-0-0	8	8	5
Mac Conkey broth	5-1-0	4-1-0	1-1-0	33	17	4
Lactose Broth	5-5-2	4-2-1	2-1-0	542	26	7
Brilliant G.L.B. Broth	5-3-1	4-0-0	2-0-0	109	19	5
Mac Conkey broth	4-0-0	2-0-0	2-0-0	13	5	5
Lactose Broth	4-0-0	1-0-0	1-0-0	13	2	2
Brilliant G.L.B. Broth	2-0-0	2-0-0	1-0-0	5	5	2
Mac Conkey broth	3-1-0	3-0-0	0-0-0	11	8	< 2
Lactose Broth	5-1-0	4-1-0	4-1-0	33	17	17
Brilliant G.L.B. Broth	3-1-0	3-1-0	3-0-0	11	11	8
Mac Conkey broth	4-3-0	4-2-0	4-2-0	27	22	22
Lactose Broth	5-0-0	5-0-0	4-0-0	23	23	13
Brilliant G.L.B. Broth	5-0-0	5-0-0	3-1-0	23	23	11
Mac Conkey broth	1-0-0	1-0-0	1-0-0	<2	< 2	42
Lactose Broth	5-0-0	5-0-0	2-0-0	23	23	5
Brilliant G.L.B. Broth	3-0-0	2-0-0	2-0-0	8	5	5
Mac Conkey broth	0-0-0	0-0-0	0-0-0	4 2	2	2
Lactose Broth	5-0-0	4-0-0	3-0-0	23	13	8
Brilliant G.L.B. Broth	5-2-0	5-2-0	5-2-0	49	49	49
Mac Conkey broth	5-0-0	5-0-0	2-0-0	23	23	5
Lactose Broth	5-0-0	1-0-0	1-0-0	23	2 2	
Brilliant G.L.B. Broth	3-0-0	1-0-0	1-0-0	23	2	2 2
lac Conkey broth	2-0-0	1~0-0	0-0-0	5	2	< 2
actose Broth	5-0-0	2-0-0	2-0-0	5	5	5
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2
lac Conkey broth	4-0-0	3-0-0	0-0-0	13	2	<2
actose Broth	5-4-0	3-2-0	3-2-0	130	14	14
Brilliant G.L.B. Broth	3-0-0	2-0-0	2-0-0	8	5	5
lac Conkey broth	5-4-1	3-3-1	3-0-0	172	21	8
actose Broth	5-1-1	5-1-0	0-0-0	46	33	<2
Brilliant G.L.B. Broth	5-3-1	3-2-1	3-0-0	109	17	8
iac Conkey broth	5-3-2	5-2-2	2-2-1	141	94	12
actose Broth	5-2-2	5-1-1	4-1-1	94	94 46	
Brilliant G.L.B. Broth	4-1-0	4-1-0	1-0-0	17	17	21 2

Tableau 4 (suite)

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>contaminé</u> Nature de l'échantillon : sédiment

27 6	Formule	des tubes por	sitifs	Nombre le	plus probabl		
Type de bouillon utilisé	Coliformes totaux	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli	_
	4-1-0	2-0-0	1-0-0	17	5	2	
fac Conkey Broth	-	3-1-0	3-0-0	33	11	11	
actose Broth	5-1-0	5-0-0	3-0-0	23	23	8	
brilliant G.L.B. Broth	5-0-0			14	14	9	
lac Conkey Broth	3-2-0	3-2-0	2-2-0		46	46	
actose Broth	5-1-0	5-1-1	5-1-1	33	22	22	
Brilliant G.L.B. Broth	4-2-0	4-2-0	4-2-0	2.2		1 10 0	
		5-1-0	3-0-0	348	33	8	
lac Conkey Broth	5-5-1	-	3-2-0	40	2.7	14	
actose Broth	4-4-2	4-3-0	3-2-0	_	_	-	
Brilliant G.L.B. Broth	-	-				7	
	5-3-2	3-1-0	2-1-0	141	11		
Mac Conkey Broth	5-3-2	4-1-0	4-1-0	141	17	17 8	
Lactose Broth	5-2-1	4-0-0	3-0-0	70	13	Ö	
Brilliant G.L.B. Broth			2-1-1	94	17	9	le l
Mac Conkey Broth	5-2-2	3-2-1		221	109	109	
Lactose Broth	5-4-2	5-3 - 1	5-3-1	94	32	2	
Brilliant G.L.B. Broth	5-2-2	4-2-2	1-0-0			1/	
THE R O SHOULD BE SEEN THE SHOULD BE	5-3-1	5-3-1	3-2-0	109	109	14 23	
Mac Conkey Broth	5-0-1	5-0-1	5-0-0	31	31		
Lactose Broth	-	2-0-0	0-0-0	22	5	< 2	
Brilliant G.L.B. Broth	4-2-0			109	5	2	8
Mac Conkey Broth	5-3-1	2-0-0	1-0-0	31	23	8	
Lactose Broth	5-0-1	5-0-0	3-0-0	22	13	13	
Brilliant G.L.B. Broth	4-2-0	3-0-2	3-0-2	22		V 4 113	1
	4-2-2	2-1-1	1-0-0	32	9	2	
Mac Conkey Broth		5-0-0	4-0-0	23	23	13	
Lactose Broth	5-0-0	J-U-U	700				
Brilliant G.L.B. Broth					27	17	
Mac Conkey Broth	4-3-0	4-3-0	3-3-0	27	141	<2	
Lactose Broth	5-3-2	5-3-2	0-0-0	141	17	42	
Brilliant G.L.B. Broth	5-1-0	4-1-0	0-0-0	33	A		
		1-0-0	1-0-0	23	2	2	
Mac Conkey Broth	5-0-0		4-0-1	31	17	17	
Lactose Broth	5-0-1	4-0-1 3-0-0	2-0-0	33	8	5	
Brilliant G.L.B. Broth	5-1-0	3-0-0				<2	
	0-0-0	0-0-0	0-0-0	< 2	< 2	5	
Mac Conkey Broth	3-0-0	3-0-0	2-0-0	8	8	<2	
Lactose Broth		0-0-0	0-0-0	< 2	<2	< 2	
Brilliant G.L.B. Broth			1-0-1	21	33	. 4	
Mac Conkey Broth	3-3-2	3-3-2		32	14	5	
Lactose Broth	4-2-2	3-2-0	2-0-0	14	8	2	
Brilliant G.L.B. Brot	h 3-1-1	3-0-0	1-0-0	14	O		

Tableau 5

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>propre</u> Nature de l'échantillon : eau de surface

Туре	Formule	des tubes po	sítifs	Nombre le	plus probabl	e, 100	
de bouillon utilisé	Coliformes totaux	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli	
Mac Conkey Broth	3-3-0	1-1-0	1-0-0	17	4	2	
Lactose Broth	4-2-0	0-1-0	0-0-0	22	2	<2	
Brilliant G.L.B. Broth	1-0-0	1-0-0	1-0-0	2	2	2	
Mac Conkey Broth	4-1-0	1-0-0	1-0-0	17	2	2	
Lactose Broth	5-1-1	1-0-0	0-0-0	36	2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey Broth	0-0-0	0-0-0	0-0-0	<2	<2	< 2	
Lactose Broth	2-0-0	0-0-0	0-0-0	5	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	<2 €2	< 2	
Mac Conkey Broth	4-1-0	2-1-0	0-0-0	17	7	< 2	
Lactose Broth	5-0-0	1-0-0	0-0-0	23	2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey Broth	3-0-0	0-0-0	0-0-0	8	<2	<2	
Lactose Broth	4-0-0	1-0-0	0-0-0	13	2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	<2	<2	
Mac Conkey Broth	1-2-0	1-2-0	1-0-0	6	6	2	-
Lactose Broth	5-2-1	3-1-0	1-0-0	70	11	2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2	< 2	<2	
Mac Conkey Broth	4-1-0	1-1-0	0-0-0	17	4	<2	
Lactose Broth	5-0-0	5-0-0	0-0-0	23	2.3	<2	
Brilliant G.L.B. Broth	4-0-0	2-0-0	0-0-0	13	5	<2	
Mac Conkey Broth	2-0-0	0-0-0	0-0-0	5	<2	<2	
Lactose Broth	1-0-0	0-0-0	0-0-0	2	<2	<2	
Brilliant G.L.B. Broth					-		
Mac Conkey Broth	1-0-0	0-0-0	0-0-0	2	<2	<2	
Lactose Broth	3-0-0	0-0-0	0-0-0	8	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	< 2	<2	<2	
Mac Conkey Broth	5-0-1	2-0-0	0-0-0	31	5	<2	
Lactose Broth	5-1-0	2-0-0	0-0-0	33	Ś	<2	
Brilliant G.L.B. Broth	1-2-1	1-2-1	1-1-0	8	8	4	
Mac Conkey Broth	4-0-0	2-0-0	0-0-0	13	5	<2	
Lactose Broth	5-0-0	2-0-0	1-0-0	23	5	2	
Brilliant G.L.B. Broth	1-0-0	0-0-0	0-0-0	2	<2	<2	
Mac Conkey Broth	4-0-0	0-0-0	0-0-0	13	<2	< 2	
Lactose Broth	4-1-0	1-0-0	1-0-0	17	2	2	
Brilliant G.L.B. Broth	1-0-0	0-0-0	0-0-0	2	<2	<2 <	

Tableau 5 (suite)

Comparaison du nombre le plus probable/100 ml des coliformes totaux, des coliformes fécaux et des <u>E. coli</u> évalué par la technique des tubes multiples appliquée en trois variantes selon le type de bouillon lactosé utilisé (Mac Conkey Broth, Lactose Broth, Brilliant Green Lactose Bile Broth)

Echantillons prélevés au point <u>propre</u> Nature de l'échantillon : sédiment

	Formule	des tubes po	sitifs	Nombre le	plus probabl	e, 100	
Type de bouillon utilisé	Coliformes totaux	Coliformes fécaux	E. coli	Coliformes totaux	Coliformes fécaux	E. coli	
	4-0-0	1-0-0	0-0-0	13	2	< 2	
lac Conkey Broth	5-1-0	1-0-0	0-0-0	33	2	< 2	
actose Broth	3-1-0 3-1 - 0	2-0-0	0-0-0	11	5	< 2	
Brilliant G.L.B. Broth				< 2	<2	< 2	
Mac Conkey Broth	0-0-0	0-0-0	0-0-0	9	<2	< 2	
actose Broth	2-1-0	0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0				_
1.0	1-0-0	1-0-0	1-0-0	2	2	2	
Mac Conkey Broth	2-0-0	2-0-0	1-0-0	5	5	2	
Lactose Broth	1-0-0	1-0-0	0-0-0	2	2	<2	
Brilliant G.L.B. Broth				₹2	4 2	< 2	
Mac Conkey Broth	0-0-0	0-0-0	0-0-0	14	<2	<2	
Lactose Broth	3-2-0	0-0-0	0-0-0	<2	<2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0				
		_	-	-	-	-	
Mac Conkey Broth	2-0-0	2-0-0	1-0-0	5	5	2	
Lactose Broth		0-0-0	0-0-0	2	<2	< 2	
Brilliant G.L.B. Broth			0-0-0	<u> </u>	< 2	< 2	
Mac Conkey Broth	0-0-0	0-0-0	1-0-0	13	2	2	
Lactose Broth	4-0-0	1-0-0	0-0-0	< 2	< 2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0					
Mac Conkey Broth	5-0-0	2-0-0	0-0-0	23	5	<2	
Lactose Broth	5-0-0	2-0-0	0-0-0	23	5	2	
Brilliant G.L.B. Broth	3-1-0	2-0-0	1-0-0	11			
CONT.	100	0-0-0	0-0-0	5	< 2	<2	10
Mac Conkey Broth	2-0-0		0-0-0	8	< 2	<2	
Lactose Broth	3-0-0	0-0-0	0-0 0	-			
Brilliant G.L.B. Broth		WI	- X		<2	< 2	
Mac Conkey Broth	3-0-0	0-0-0	0-0-0	8	<2	<2	
Lactose Broth	3-0-0	0-0-0	0-0-0	8	<2 <2	<2	
Brilliant G.L.B. Broth	0-0-0	0-0-0	0-0-0	<2		A CASA SAN SAN	100
	5-3-1	2-2-0	0-0-0	109	9	<2	
Mac Conkey Broth		4-0-0	0-0-0	13	13	<2	
Lactose Broth	4-0-0 3-2-0	0-1-0	0-0-0	14	2	<2	
Brilliant G.L.B. Broth	1 3-2-0		1 2 2 2 2 2	<u> </u>	2	<2	
Mac Conkey Broth	-	1-0-0	0-0-0	<u> </u>	<2	<2	
Lactose Broth	_	2-0-0	0-0-0		<2	<2	
Brilliant G.L.B. Broth	h -	0-0-0	0-0-0	7 7		10.51	
Acidos .	0-0-0	0-0-0	0-0-0	42	<2	<2	
Mac Conkey Broth		0-0-0	0-0-0	<2	<2	<2	
Lactose Broth	0-0-0	0-0-0	0-0-0	< 2	<2	< 2	
Brilliant G.L.B. Brot	h 0-0-0	0-0-0	0 0 0				

Tableau 6

Comparaison de la sensibilité (exprimée en pourcentage) de la technique des tubes multiples selon le type de bouillon lactosé utilisé en ce que concerne la détermination des nombres les plus probables de coliformes

Type de		e plus pro mes totaux			plus prob. es fécaux/			plus proba oli/100 ml	ble des
bouillon			6	Degr	é de sensi	bilité		<u>-</u> -	
lactosé	élevé	moyen	limité	élevé	moyen	limité	élevé	moyen	limité
Lactose Broth	79,41	14,71	5,88	71,01	24,63	2,89	76,81	20,27	2,89
Mac Conkey Broth	41,17	47,11	11,76	49,27	28,98	21,73	46,37	33,33	20,28
Brilliant G.L. Bile Broth	19,11	45,58	35,29	30,43	47,82	21,73	40,57	43,47	15,98

Tableau 7

Comparaison du degré de sensibilité (exprimé en pourcentage) des trois bouillons lactosés dans la détermination de la concentration des <u>E. coli</u> dans les échantillons analysés

Type de bouillon utilisé	NPP - coliformes totaux/100 ml	NPP - coliformes fécaux/100 ml	NPP - E. coli/ 100 ml
Lactose Broth	79,41	71,01	76,81
Mac Conkey Broth	41,17	49,27	46,37
Brilliant G.L. Bile Broth	19,11	30,43	40,57

Tableau 8

Comparaison des résultats obtenus par la méthole des tubes multiples et par la membrane filtrante concernant le NPP/100 ml des E. coli

	PROP	RE				CONTA	MINEE				POLLUE	3		
Numéro des échan-	Nombre des colonies		P/100 sur		Numéro des échan-	Nombre des colonies		/100 sur		Numéro des échan-	Nombre des colonies à 44°C		P/100 sur	
tillons			B*	M*	tillons	à 44 °C 100 ml	Ľ*	В*	Μ*	tillons	100 ml	Ľ*	В*	Μ*
					11	EAUX DE S	URFACE							
5	1	<2	2	2	3	1	5	5	< 2	1	4	109	< 2	79
11	_	<2	< 2	2	9	3	7	5	4	7	3	79	14	79
17	0	<2	< 2	< 2	15	0	2	2	5	13	1	46	33	33
23	2	<2	۷2	<2	21	-	17	8	42	19	7	240	1600>	
29	0	4 2	< 2	< 2	27	7	13	11	22	25	35	46	14	9
35	0	2	<2	2	33	2	5	5	2	31	10	17	17	33
41	6	_ < 2	< 2	<2	39	22	8	49	< 2	37	-	141	426	2400
47	0	<2	<2	<2	45	26	2	2	5	43	52	63	4	<:
53	0	<2	< 2	<2	51	0	5	<2	< 2	49	40	>2400	48	240
59	2	<2	4	<2	57	12	14	5	<2	55	40	7	11	
65	0	2	< 2	< 2	63	12	<2	8	8	61	-	172	26	10
71	5	2	< <u>2</u>	< <u>2</u>	<u>69</u>	Ξ.,	21	2	12	<u>67</u>	20	14	<u>49</u>	3
						SEDIM	ENTS							
,	0	< 2	< 2	<2	4	0	11	8	2.	2	0	141	< 2	
6	2	<2	\ 2	<2	10	-	46	22	9	8	20	240	141	2
12	1	2	<2	2	16	18	14		8	14	1.50	130	< 2	2
18	1	< 2	<2	<2	22	56	17	8	7	20	5	8	13	
24	0	2	<2	-	28	40	109	2	9	26	_	<2	<2	<
30	0	2	< 2	۷2	34	8	23	<2	14	32	-	< 2	< 2	2 4
36	20	< 2	2	<2	40	140 -	8	13	2	38	= = -	<2	< 2	2 <
42 48	4	< 2	<2	<2	46	0	13	<2	2	44	-	4	< 2	2 <
	2	<2	<2	<2	52	20	<2	<2	17	50	_	8	3 <2	2 <
54 60	4	<2	< 2	<2	58	-	17	. 5	2	56	_	42	2 <2	2 <
	-	<2 <2	<2	<2	64	-	5	<2	<2	62	260	24	+ 6	5 240
66 72	_	<2	\ 2	<2	70	_	5	2	. 4	. 68	0	21		2

^{*} L = Lactose Broth

B = Brilliant Green Lactose Bile Broth

M = Mac Conkey Broth

Tableau 9

Comparaison de la sensibilité des deux techniques (tubes multiples et membranes filtrantes exprimée en pourcentage par rapport aux nombres des échantillons analysés)

Point de prélèvement	Technique tubes multiples supérieure à celle de la membrane filtrante	Technique membrane filtrante supérieure à celle des tubes multiples	Egalité
Propre	42,8	33,33	23,8
Contaminé	70,58	29,41	0
Pollué	87,5	12,5	0
Pollué	87,5	12,5	0

Annex 8

A CONTRIBUTION TO THE COMPARATIVE STUDY OF ANALYTICAL METHODS USED FOR DETECTION OF MAJOR BACTERIAL INDICATORS OF POLLUTION CAUSED BY FAECAL SEWAGE

> by D. Fuks Rudjer Boskovic Institute, Rovinj, Yugoslavia

This study is designed to contribute to the comparative study of analytical methods used for detection of the major bacterial indicators of pollution caused by faecal sewage.

Areas studied

Three sampling points were selected in the area of Rovinj, a small town of 10 000 inhabitants, situated on the West Istrian coast of the North Adriatic, with facilities for accommodating up to 40 000 tourists:

- station no. 1: near an island not under the direct influence of domestic wastewater;
- a beach close to the town harbour, where the influence of sewage discharge in
 - the vicinity is noticeable;
- station no. 3: in the harbour itself, where most of the sewage outfalls discharge into the

Methodology

2.1 Sampling methods

Seawater samples were taken with a sterile glass sampler (using an extension arm) 10 m from the coastline, and 20-50 cm below the surface. Sediment samples were taken using a Pfleger corer. The surface layer of the sample was transferred by a spatula to a sterile Petri dish. Samples were transported to the laboratory in a cooling container. The interval between sample collection and processing was never more than three hours.

The following parameters were analysed: total coliforms, faecal coliforms, faecal streptococci and BOD5 in water, and faecal coliforms in sediments.

2.2 Analytical methods

Total coliforms: the membrane filtration (MF) technique was used as described in the relevant WHO/UNEP reference method (1). The most probable number (MPN) technique was used, as described in the WHO/UNEP Guidelines for health-related monitoring of coastal water quality (2). In the former method, samples were incubated for 24 hours at 36°C±1°C using Mac Conkey Broth. Gas and acid production were identified and used for the MPN calculation.

Faecal coliforms: the same methods as described above for total coliforms were used. In the MF technique, samples were incubated for 24 hours on solid M-FC-agar at 44.5°C±0.2°C, blue colonies being counted. In the MPN technique, samples were incubated for 24 hours at 36°C±1°C using Mac Conkey Broth. Positive tubes were tested by transfer to individual Mac Conkey tubes and incubated at 44.5°C±0.2°C for 24 hours. Confirmation was performed using the indole production test, acid, gas and indole production in the tubes being used for the MPN calculation.

Faecal streptococci: the membrane filtration technique as described in the WHO/UNEP Guidelines (2) was used, while the MPN method was used as described in the APHA Standard Methods (3). In the former, samples were incubated for 48±3 hours at 44.5°C±0.2°C on KF-streptococcus-agar, dark red colonies being counted. In the latter, samples were incubated for 48 hours at 36°C±1°C on Azide-Dextrose Broth. Due to lack of Ethyl-Violet-Azide Broth, the confirmatory test was performed using a Bromocresol-purple Azide Broth. Incubation was performed at 36°C±1°C for 48 hours. Increasing turbidity and colour change to yellow were used for MPN calculation. Bromcresol-purple Azide Broth (g/1) was prepared as follows: peptone from casein - 10.00; yeast extract - 10.00; D-glucose - 5.0; NaCl - 5.0; K2HPO₄ - 2.7; KH2PO₄ - 2.7; NaN - 0.5; Reconstant purple - 0.032 2.7; NaN3 - 0.5; Bromcresol- purple - 0.032.

8005: the technique described in the APHA Standard Methods (3) was used for determination of the biochemical oxygen demand. Samples were incubated in 250 ml bottles.

Faecal coliforms in sediment: the MPN technique described above for determination of faecal coliforms in seawater was used for this parameter.

Results and discussion

The results obtained during the experimental period are presented numerically in Table 1 and graphically in Figures I to 3. Analysis of the data shows significant differences in the level of pollution between the three stations (Table 1). The concentration of bacterial indicators as well as the biochemical oxygen demand value confirm that station no. 1 is located in an unpolluted area, station no. 2 is subject to wastewater pollution, and no. 3 is polluted by discharges from sewage outfalls. Although the values of $80D_5$ obtained showed differences in pollution levels between stations, correlation between the BOD_5 values and the concentration of bacterial indicators was obtained only at station no. 3 (r = 0.87-89). The number of faecal coliforms recorded in sediments confirmed the influence of location of sewage outfalls on the quality of the seawater in the various stations.

Figures 1-3 show some discrepancy between the results obtained using the MF and MPN techniques respectively. Higher values of bacteria were obtained in 67% of samples determined by the MF technique, as compared to those obtained using the MPN method. Extreme differences were noticed for faecal streptococci at station no. 2 where in 5 out of 6 samples, MF values were out of the MPN 95% confidence limits. Statistical evaluation of the results showed significant differences between MF values and MPN values only in the case of faecal streptococci at station no. 2 (P = 0.001).

The phenomenon whereby the membrane filter technique produces lower counts than the multiple tube test or most probably number (MPN) techniques in the bacteriological examination of chlorinated wastewaters has not previously been recorded in the coastal waters of Royinj.

In comparing the relative merits of the MF and MPN techniques, the precision obtained with a single membrane filter was five times greater than that of a five-tube MPN (4). Also, the MF method gave much better replication (5). Results obtained in the present experiment confirmed previous results recorded in MF/MPN comparisons in the analysis of coastal waters from the Rovinj area, although the media used were different (6).

Although the MF and MPN tests for the enumeration of total and faecal coliforms as well as for that of faecal streptococci do not always yield exactly the same results, it is clear that both methods do supply substantially the same information.

In summary, membrane filtration using M-FC-agar provides a direct bacterial count and requires only 24 hours to complete. A commercial product which automatically switches incubation temperature is available to make this procedure more practical for routine application.

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Table 1

The variation of parameters covered by methodology testing exercises

Station No.	date of sampling	hour of sampling	Total o	Total coliforms MF MPN	Faecal of MF n/	Faecal coliforms MF MPN n/100 m1	Faecal MF	Faecal streptococci	MPN	coliforms in sediments n/l g
	20.05.82	10.00	6	0	2	0	0		5 5	0
	4.06.82	08.45	0	0	0	0	0 9		o c	, ,
	17,06,82	00.60	20	2	16	0 (71		o C	0
	22.06.82	00.10	3	2		7 6	→ <		ο α	6
	14.07.82	12.45	೯	∞	1	0 (0 -) C	4
	29.07.82	00.60	22	2	ıΩ	5	# 1a		>	
				C	300	204	200		110	2
	20.05.82	06.30	280	350	202		420		4	0
	4.06.82	09.25	800	920	676	0 0	000		13	7
	17,06,82	09.20	610	240		240	076		7.6	14
	22.06.82	10.50	1 050	140	1 015	140	790		, ,	۲۰
	14.07.82	13,15	70	42	55	E E	90		. 011	6
	29.07.82	04.60	2 600	240	400	350	619		011	ı *
		6	0	5 400	3 300	3 500	6.650	7	200	33
3	20.05.82	10,30	3 400	000 71			13 600	3	3 500	67
	4.06.82	09.40	20 000	14 000	21 500	24 000	47 200	7	7 900	220
	17.06.82	10.30	87 500			000 68		54	000	110
	22.06.82	10.10	135 000	92 000	1 000				09	22
	14.07.82	13.20	050			007	1 250	3	700	240
	29 07.82	09.45	2 700	3 500	1 700	490)	

Figure 1

The variation of total coliforms, faecal coliforms and faecal streptococci in six water samples taken at Station No. 1*

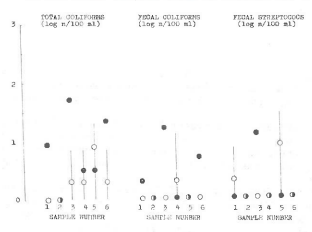


Figure 2

The variation of total coliforms, faecal coliforms and faecal streptococci in six water samples taken at Station No. 2*

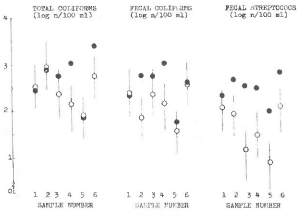
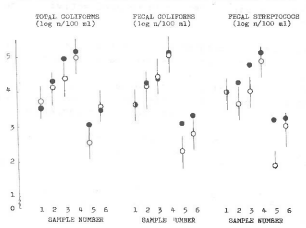


Figure 3

The variation of total coliforms, faecal coliforms and faecal streptococci in six water samples taken at Station No. 3*



*Results of the MF method indicated by (o), results of the MPN method indicated by (o) with 95% confidence limit.

Annex 9

REPORT ON THE JOINT INTERCALIBRATION EXERCISE ON MICROBIOLOGICAL METHODS

Introduction

During the course of the pilot phase of the Joint Coordinated Mediterranean Pollution Monitoring and Research Programme (MED POL Phase II), draft reference methods were produced for use in the long-term monitoring programme. These methods included the following:

- determination of total coliforms in seawater by the membrane filtration culture method;
- determination of faecal coliforms in seawater by the membrane filtration culture method; determination of faecal streptococci in seawater by the membrane filtration culture method;
- determination of faecal coliforms in bivalves by the multiple test-tube method.

The above four methods were developed by UNEP's Regional Seas Programme Activity Centre (RS/PAC) in cooperation with the WHO Regional Office for Europe. By mid-1982, all four had reached a sufficient stage of development to be tested by individual laboratories, following which an intercalibration exercise could be organized prior to final review.

Organization 2.

A selected number of Mediterranean institutions were invited to test the reference methods under their local environmental conditions during the late summer/early autumn of 1982. They were also invited to send representatives to participate in a joint intercalibration exercise in Rome on 22-23 November 1982, and in a consultation meeting to review the methods, also in Rome, from 24 to 26 November 1982.

The following laboratories were invited:

- Centre de Recherche océanographique et des Pêches, Alger, Algeria;
- Cellule de Lutte contre les Pollutions marines, Service maritime des Bouches-du-Rhône,
- Marseille, France;
- Environmental Pollution Control Project, Athens, Greece;
- Environmental Health Laboratory, Hadassah Medical School, Jerusalem, Israel;
- Institute of Water Supply and Wastes Disposal, University of Naples, Italy;
- Public Health Laboratory, Department of Health and Environment, Valletta, Malta;
- Institut national d'Hygiène, Ministère de la Santé publique, Rabat, Morocco;
- Jefatura Provincial de Sanidad, Malaga, Spain;
- Environmental Engineering Department, Middle East Technical University, Ankara, Turkey;
- Institute of Biology, University of Ljubljana, Portoroz, Yugoslavia.

Selection of laboratories was made on the basis of (a) participation in MED POL Phase I (MED VII - Coastal Water Quality Control), (b) intended participation in the long-term phase of MED POL, (c) geographical distribution. Because of financial limitations, account was also taken of invitations already issued for participation in related activities during 1982, and plans for similar exercises during 1983.

Nine Mediterranean laboratories accepted the invitation to test the methods and sent representatives to participate in the joint intercalibration exercise and consultation meeting.

By arrangement with UNEP's Regional Seas Programme Activity Centre, the three following laboratories outside the Mediterranean region were also invited and accepted to participate in the testing and send representatives to the joint intercalibnfation exercise and consultation meeting:

- Environment Pollution Division, Ministry of Health, Sha'ab, Kuwait;
- Bacteriological Laboratory, Ministry of Health, Oman;
- The Caribbean Environmental Health Institute, St Lucia, West Indies.

Also by arrangement with UNEP's Regional Seas Programme Activity Centre, the following experts were invited to attend the joint intercalibration exercise and the consultation meeting in an advisory capacity: Dr G.I. Barrow (United Kingdom), Dr M. Bernhard (Italy), Professor E. Geldreich (USA) and Mr K. Krongaard Kristensen (Denmark).

Copies of the reference methods were transmitted to the laboratories for testing during late August/early September 1982.

Proceedings and results

The joint intercalibration exercise was held in the laboratories of the Istituto Superiore di Sanità, Rome, from 22 to 25 November 1982. A list of participants is appended. On the last two days (24 and 25 November), activities were limited to analysis of results, and the time-table arranged for cohesion with the consultation meeting.

The exercise was limited to the first three reference methods, i.e.:

- determination of total coliforms in seawater by the membrane filtration culture method;
- determination of faecal coliforms in seawater by the membrane filtration culture method;
- determination of faecal streptococci in seawater by the membrane filtration culture method.

It was originally intended that, apart from determination of the three parameters (total coliforms, faecal coliforms and faecal streptococcci) by the membrane filtration (MF) method, participants could also carry out parallel determinations on the same samples, utilizing the most probable number (MPN) method. Detailed instructions were therefore prepared in this regard for the following samples:

- No. 1 effluent (fresh water) MF and MPN
- No. 2 "clean" area (seawater) MF and MPN
- No. 3 intermediate area (seawater under the possible influence of a pollution source from which sample No.1 was taken) - MF and MPN
- No. 4 sand from sampling point No. 2 MPN
- No. 5 sand from sampling point No. 3 MPN.

During the exercise, considering the available time, it was decided to limit work to determination of the three parameters by the MF method only (i.e. as detailed in the draft reference methods). Only samples Nos 1, 2 and 3 were therefore utilized.

Participants were divided into six teams of two persons each. Results obtained by individual teams were more or less in agreement, although differences were statistically significant, even if slight, in some cases. A preliminary analysis of part of the data was also carried out.

The original instructions as given to participants, the results obtained and the preliminary statistical analysis, are reproduced herewith.

Conclusions

Considering that this was the first laboratory exercise of this nature organized within the framework of MED POL, the results can be described as satisfactory, and have provided the basis on which further intercalibration exercises both at national and regional levels can be organized in future years. In addition, the occasion provided the opportunity for participants to compare techniques during actual work-performance, rather than through discussion only.

Instructions given to participants in intercalibration exercise on microbiological reference methods

SAMPLES 1.

No. 1 = Effluent (fresh water).

No. 2 = Clean area (sea water).

No. 3 = Intermediate area (sea water under the possible influence of the pollution source No. 1).

No. 4 = Sand of sampling point No. 2.

No. 5 = Sand of sampling point No. 3.

SAMPLING METHOD 2.

The samples are collected directly by hand, in sterilized containers, and are carried out by cooling containers (4-10°C). Storage and transportation time = maximum 24 hours.

ANALYTICAL METHODS

WATER SAMPLES (No. 1, 2 and 3) 3.1

TOTAL COLIFORMS 3.1.1

Membrane filter method 3.1.1.a

M-Endo-Agar 3.1.1.a.1

Filter 100 ml of sample (or its dilution). Incubate 24 hours at 36±1°C on solid M-Endo-Agar. After incubation, read for typical colonies (pink or red). Calculate for bacterial density (see Form A).

M-FC-Broth 3.1.1.a.2

Filter and incubate as in 3.1.1.a.1 on M-FC-Broth. After incubation, read for typical blue colonies. Calculate for bacterial density (see Form A).

Most probable number method (MPN) 3.1.1.b

Inoculate Lactose Broth tubes with the sample or its dilutions. Incubate 18-24 hours at 36±1°C.

Read for gas production. For negative tubes, incubation is continued for another 24 hours (total incubation time = 48±3 hours).

Positive tubes are tested by transferring a small inoculum from each positive tube to individual Brilliant Green Lactose Bile Broth tubes and incubating at 36±1°C for 48 hours.

Read for gas production. Calculate for MPN (see Form A).

FAECAL COLIFORMS 3.1.2

Membrane filter method 3.1.2.a

M-FC-Agar 3.1.2.a.1

Filter as in 3.1.1.a.1.

Incubate 24 hours on solid M-FC-Agar at 44±0.2°C (water-bath). After incubation, read for blue colonies. Calculate for bacterial density (see Form A).

M-FC-Broth 3.1.2.a.2

Filter as in 3.1.1.a.l.

Incubate 24 hours on M-FC-Broth at 44.5±0.2°C (water-bath). After incubation, read for blue colonies. Calculate for bacterial density (see Form A).

3.1.2.b Most Probable Number Method (MPN)

Inoculate tubes, incubate and read for gas production, as in 3.1.1.b. Positive tubes are tested by transferring a small inoculum of each positive tube to individual EC-Broth tubes and incubating at 44.5±0.2°C for 24 hours in a water-bath. Calculate for MPN (see Form A).

3.1.3 FAECAL STREPTOCOCCI

3.1.3.a Membrane filter method

3.1.3.a.1 KF-Agar (at 44°C)

Filter as in 3.1.1.a.1. Incubate 48 ± 3 hours at $44\pm0.2\,^{\circ}\text{C}$ on KF-Agar (water-bath). After incubation read for red colonies or colonies with a red center. Calculate for bacterial density (see Form B).

3.1.3.a.2 KF-Agar (at 36°C)

Filter as in 3.1.1.a.1. Incubate 48±3 hours at 36±1°C on KF Agar. After incubation read for red colonies or colonies with a red center. Calculate for bacterial density (see Form B).

3.1.3.a.3 M-Enterococcus Agar

Filter as in 3.1.1.a.1. Incubate 48 hours at 36±1°C. After incubation read for pink or red colonies. Calculate for bacterial density (see Form B).

3.1.3.b.1 Most Probable Number Method (MPN) at 36°C

Inoculate tubes of Azide Dextrose Broth with the sample or its dilutions. Incubate at $36\pm1\,^{\circ}\text{C}$ for 24+24 hours. Read for turbidity. Positive tubes have to be confirmed on EVA Broth (24+24 hours at $36\pm1\,^{\circ}\text{C}$). Read for turbidity and formation of a "purple" bottom. Calculate for MPN (see Form B).

3.1.3.b.2 Most Probable Number Method (MPN) at 44°C

Inoculate tubes of Azide Dextrose Broth with the sample or its dilutions. Incubate at 44±0.2°C for 24+24 hours (water-bath). Read for turbidity. Positive tubes have to be confirmed on EVA Broth (24+24 hours at 44±0.2°C in water-bath.) Read for turbidity and formation of "purple" bottom. Calculate for MPN (see Form B).

ITEMS AVAILABLE TO EACH PARTICIPANT

1.	Samples
1.1	Samples of water No. 3
1.2	Samples of sand No. 2
2.	Membrane filters (pores of 0.45 micron, sterile, ready for use), No. 20
3.	Pipets (sterile)
3.1	Graduated pipets ml 10 No. 10
3.2	Graduated pipets ml 5 No. 10
3.3	Graduated pipets ml 1 No. 10
3.4	Ungraduated (Pasteur) No. 10
4.	Petri dishes (sterile) No. 10
5.	Dílution tubes (sterile, containing ml 9 of phosphate buffer, pH 7.2) No. 40
6.	Culture media
6.1	Lactose Broth
6.1.1	Lactose Broth (2 X) Durham tubes containing ml 10, No. 40
6.1.2	Lactose Broth, Durham tubes (1 X) containing about m1 10, No. 40
6.2	Brilliant Green Lactose Bile Broth, Durham tubes containing about ml 10, No. 60
6.3	EC Broth, Durham tubes, containing about ml 10, No. 60
6.4	Azide Dextrose Broth
6.4.1	Azide Dextrose Broth (2 X), tubes containing about ml 10, No. 40
6.4.2	Azide Dextrose Broth (1 X), tubes containing about ml 10, No. 40
6.5	Ethyl Violet Azide (EVA) Broth, tubes containing about ml 10, No. 60
6.6	M-Endo Agar, in Petri dishes, ready for use, No. 10
6.7	M-FC Agar, in Petri dishes, ready for use, No. 10
6.8	KF Agar, in Petri dishes, ready for use, No. 10
7.	Racks for tubes
8.	Cas flame
9.	Wire loop

Glass marker

10.

Form	A	(Wate	r)
LOLIII	2 6	(110000	/

		For	m A (Water	r)						
	Country Somple is	; ;									
Leboratory	50 Country St. 44.5°C Country St. 50 Web.		FC MPN/100 mg								
Portecipating behoratory	M-Endo-ogartul M-FC Broth 44.5°C M. Fc Age 44.5°C Country	ठा फिली.	T.C MPN/100 W/		. 2014				-		
	36°C M. Fo Agn	be Jest me	EC II								
. Mf metho	M.Fc Broth	Mulexipe. Tu	BI EC I								
Total and Pecel coliforus. MF method	- Ondo og + (4)	colipuns - Multiple Tube Jest method.	B4LBB 84L8811								
e and pec	2 2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	pecal	L.B.								
Tota	00 100 100 100 100 100 100 100 100 100	Total and	INDENTINGUAGION		678	527	748	48	48	77.4	7 80
		_			7007	7 -07	10.1	10.3	r 5.01	10.5 4	r 3.01

Form B (Water)

M.Ewderac, Agar 36°C Country Sample no	FECAL STREPTOCOCCI. MAN METHOD	ENABroth II FS MPN/1020 /								
F-AGAR 36°C	Pr. Let 36°C	Az. Dext. Br. EVA broth I								
A Cuellum Lethol		incentum incubation	10 ml 24	787 2	1 4 24	1 u 48	7 x 7	4 u 24	1 u 124	7 24
diention (10° 10° 10° 10° 10° 10° 10° 10° 10° 10°		elibrion in	001	-	70-1	70-5-01	70.3	70-4	-	9_01

Form C (Sand-Sediment)

Patticitating Laboratory		Country	Sample 40
1-	1	9	4,

Feese coliorms- Multiple tube test method	insculum ml L.B. E.C. I E.C. II FC MPN	10 kg	1 2 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	77 /	1 57 Y	7 78		24	λ μη γ
Fecal	dilution	, or	700	70~1	70-7	€-0)	h-01	5-0)	40-6

SUMMARY RESULTS OF INTERCALIBRATION EXERCISE

Sample No.	Sample Test Amount	Team 1	Results Team 2	obtained (med	ian and range) Team 4	Team 5	Team 6
Faecal	Coliforms 0.1 ml	190(156-210)	190(151-224)	110(102-118)	152(148-186)	157(138-183)	00
1	0.01 ml	15 (10-22)	18 (17-26)	22 (17-54)	20 (0-24)	28 (27-29)	30 (24-32)
2	100 ml	1 (1-2)	5 (4-8)	2 (1-3)	4 (2-4)	5 (5-5)	5 (2-6)
2	100 m1	1 (0-1)	0 (0-1)	0 (0-0)	0 (0-0)	1 (1-2)	0 (0-0)
3	100 m1	11 (9-12)	11 (9-12)	18 (12-19)	6 (6-6)	10 (8-15)	10 (8-14)
3	10 ml	3 (2-3)	3 (2-3)	1 (0-1)	3 (0-5)	1 (0-1)	3 (0-6)
Faecal	Streptococ	ci					
1	1 m1	67 (65-72)	47 (41-52)	60 (55-83)	67 (62-74)	52 (51-53)	51 (50-56)
1	0.1 ml	6 (3-8)	3 (2-5)	9 (4-10)	6 (5-7)	5 (2-6)	7 (7-8)
1	0.01 ml	1 (0-1)	-	0 (0-1)	1 (0-2)	1 (0-3)	0 (0-1)
2	100 ml	7 (1-8)	5 (2-6)	7 (2-8)	5 (4-5)	5 (4-7)	5 (3-7)
2	10 ml	0 (0-1)		0 (0-0)	1 (0-1)	1 (0-1)	0 (-1)
3	100 ml	10 (7-13)	12 (4-16)	11 (7-17)	11 (10-11)	13 (6-14)	13 (12-24)
3	10 m1	3 (2-3)	-	1 (1-1)	1 (0-1)	1 (0-3)	1 (0-2)

INTERCALIBRATION EXERCISE: PRELIMINARY ANALYSIS

The analysis has been limited to the first collaborative experiment, concerning water samples at 0.1 and 0.01 dilution factor.

The following hypotheses have been assumed:

- A Poissonian statistical distribution of values,
- An approximate Gaussian behaviour of such distribution in the range of data examined.

Under these conditions, confidence limits of detected values have been computed, as shown in Fig. 1.

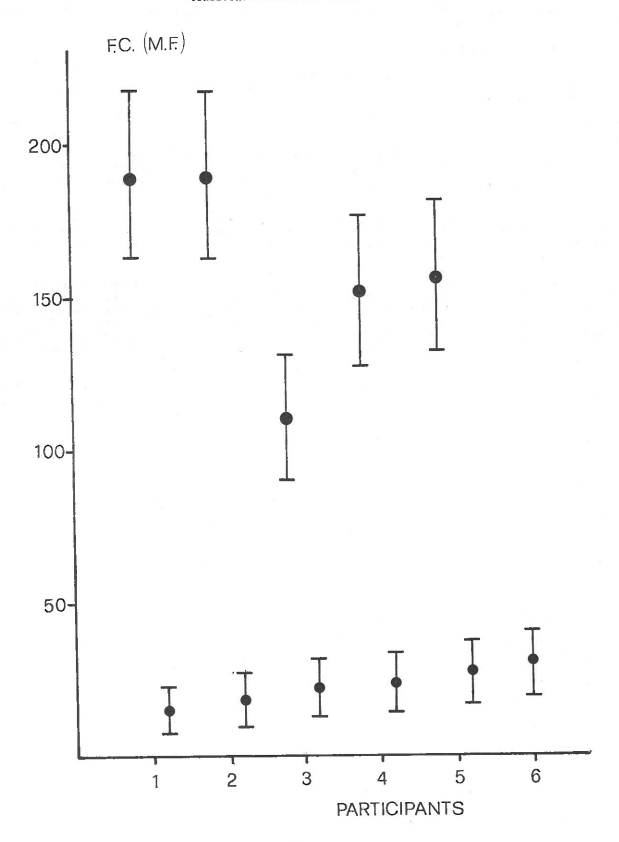
The expected values of the measured parameter, as estimated on the basis of the two determinations available for each participant (0.1 and 0.01 dilution factor samples), are reported in Fig. 2, together with the corresponding confidence limits. The width of last confidence interval is due to the lack of the 0.1 dilution datum.

The agreement among the estimates appears to be good. However, the hypothesis of some slight difference "among participants", statistically significant, may be posed. Moreover, at least in one case, a slight significant difference may be hypothesized between the two estimates carried out at the two different dilution factors.

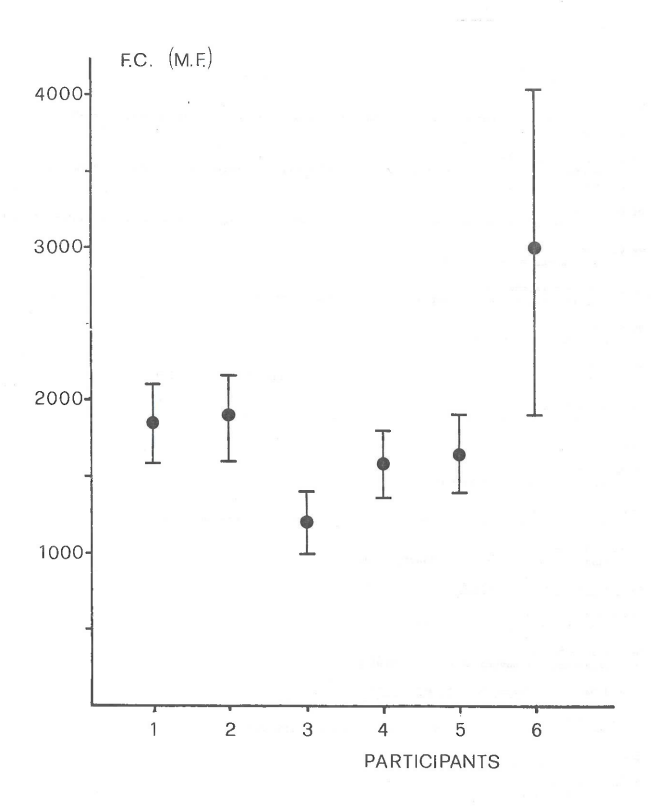
In any case, these nearly negligible differences may be considered as largely acceptable, if considered in the light of practical needs in environmental monitoring. If this reasonable point of view is assumed, the experiment outcomes appear to be fairly satisfying. Analogous conclusions may be expected for the two other series of measurements. Due to the little numbers of F.C. counted, the Gaussian approximation of Poissonian statistical distribution appears not acceptable in this case. Therefore, a reliable data evaluation requires much more computer work, and a simple estimate is not possible.

In any case, all the hypotheses here presented need to be verified with further study and have to be considered as a rough preliminary estimate. The comments of participants are necessary to design a complete analysis of data.

 $\frac{\text{Figure 1}}{\text{Confidence limits of detected values}}$



 $\frac{\text{Figure 2}}{\text{Expected values and confidence limits of measured parameters}}$



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Annex 10

OUTLINE SUMMARIES OF REFERENCE METHODS, AS MODIFIED

Summary A: total coliforms in sea water (Reference Method No. 2)

Method:

membrane filtration

Medium:

M-endo agar (not to be autoclaved)

Incubation temperature:

36±1°C (air incubator)

Time:

24 hours

Colour of colonies:

Pink to red with golden sheen

Confirmation:

AG on Mac Conkey broth 48 hours at $36\pm1^{\circ}\text{C}$, or G on brilliant green broth 48 hours at $36\pm1^{\circ}\text{C}$ (not systematic, carried out

only initially and thereafter when in doubt)

Summary B: faecal coliforms in sea water (Reference Method No. 3

Method:

membrane filtration

Medium:

 $\ensuremath{\mathsf{m-FC}}$ agar (not to be autoclaved) add rosolic acid only if necessary

Incubation temperature:

44.5±0.2°C (water-bath)

Time:

24 hours

Count:

blue colonies

Confirmation:

AG on Mac Conkey broth 24 hours at 44.5°C, or G on brilliant green broth 24 hours at 44.5°C, and Indole + in tryptone water 24 hours at 44.5°C (not systematic, carried out only

initially and thereafter when in doubt)

Summary C: faecal streptococci in sea water (Reference Method No. 4)

Method:

membrane filtration

Medium:

KF agar (not to be autoclaved)

Incubation temperature:

36±1°C (air incubator)

Time:

48 hours

Colonies:

pink to maroon

Confirmation:

not necessary

Summary D: faecal coliforms in bivalves (Reference Method No. 5)

Method:

MPN

Medium:

lactose broth (Presumptive Test) at 36±1°C (48 hours)

Mac Conkey broth at 44.5±0.2C (24 hours

or brilliant green broth at 44.5±0.2°C (24 hours and Indole

test at 44.5±0.2°C (24 hours)

Annex 11

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