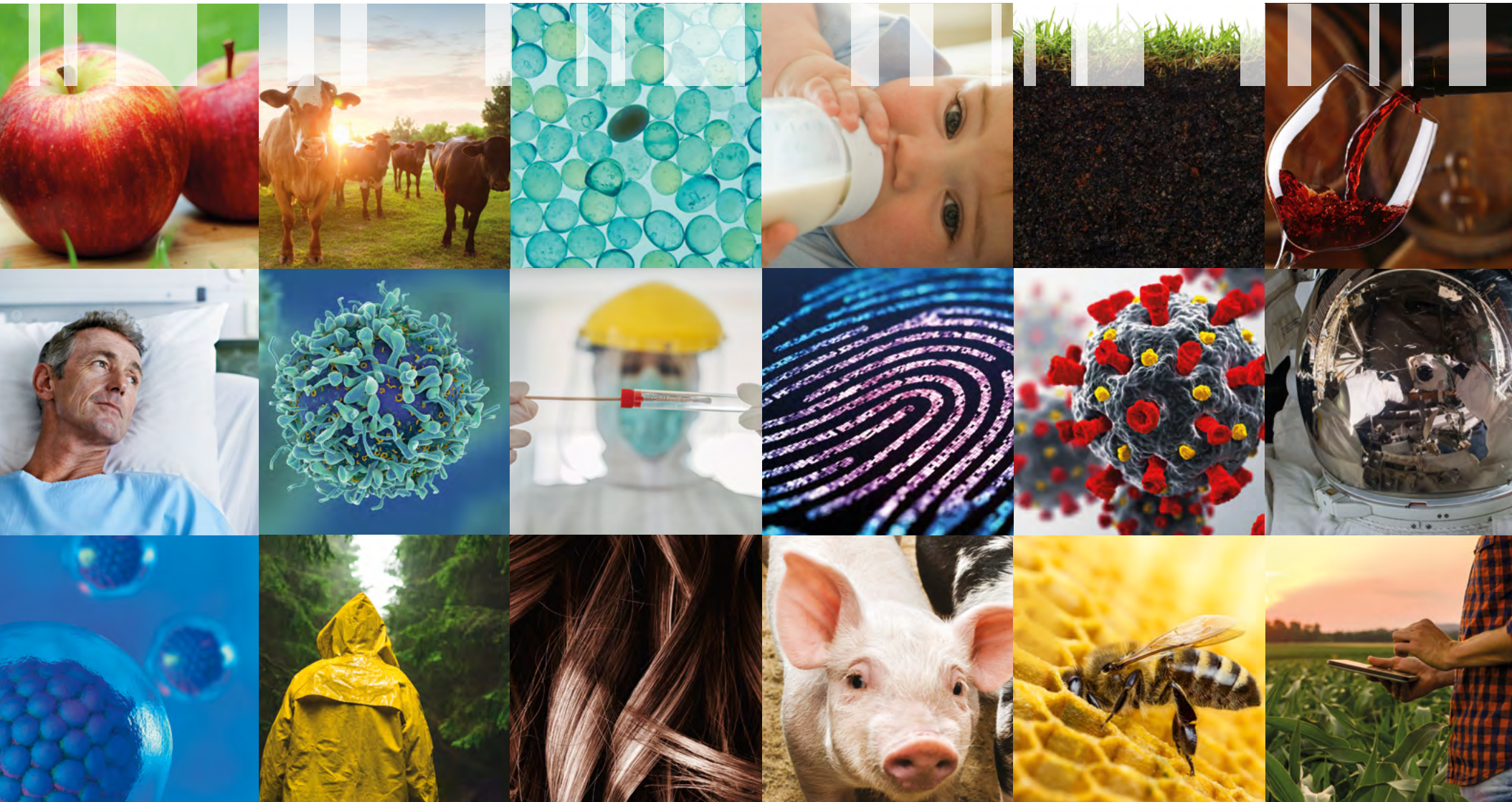


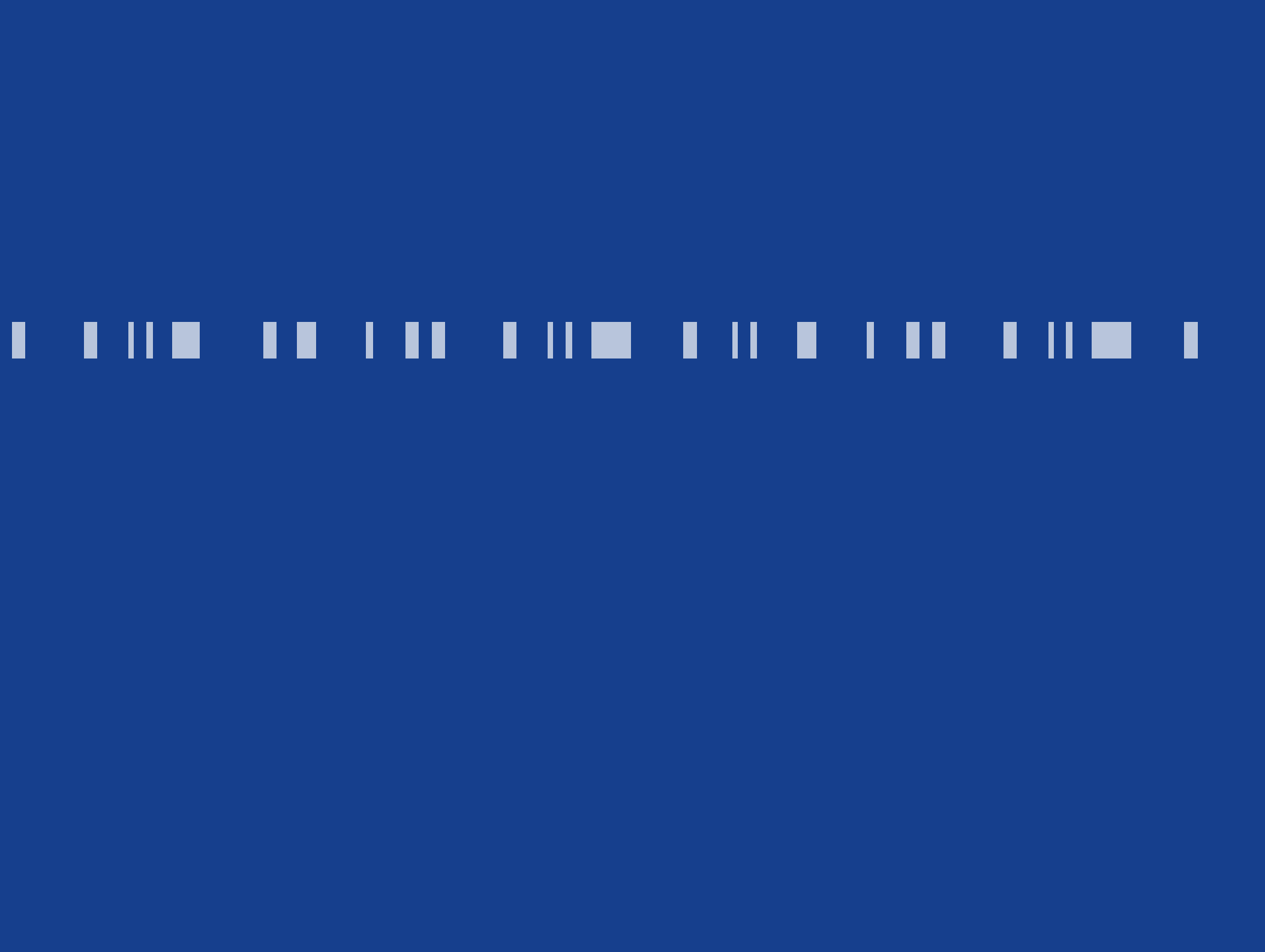
Science that makes an impact

The stories behind some of our innovations

JUNE 2025

 **eurofins**
Testing for Life





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EUROFINS LABORATORIES HAVE REGULARLY BEEN THE FIRST IN THE WORLD TO DEVELOP NEW TESTING TECHNOLOGIES THAT HAVE HAD AN IMPACT ON LIFE

OVER 35 YEARS 950+ LABORATORIES CA. 63,000 EMPLOYEES

33 MILLION TESTS PERFORMED EVERY MONTH.
WE'RE HERE TO KEEP OUR WORLD SAFE.

Since its foundation in 1987, Eurofins has contributed significantly to the advancement of science in testing for life. We are proud of the discoveries and advancements our specialists have made in myriad sectors, ranging from food to pharma to forensics testing.

Our laboratories around the world are helping to improve cancer treatments, catch criminals, safeguard the health of bees, ensure our food and water is safe, determine paternity, and beat viruses like COVID-19. Underpinning our work in these areas and more, Eurofins has one of the world's largest and most varied databases, rich with fingerprints of thousands upon thousands of foodstuffs and drugs.

The complexity of our analyses, often searching for the minutest traces of a substance, mean our activities in these areas necessarily sometimes take years of painstaking research and ongoing improvements of our numerous methods. In other areas, we have responded overnight to global health crises.

Wherever we are and however long it may take, our work helps to make life safer.

This booklet presents a mere fraction of the hundreds of stories from across our activities about our scientists' great work to push the frontiers of Testing for Life.

PROTECTING TRANSPLANT PATIENTS AND

SAVING LIVES

Immunity at the heart of the matter

Surgeons worldwide perform more than 150,000 solid organ transplants annually, with successful transplantation of hearts, lungs, livers and kidneys, saving and changing thousands of lives. After complex surgery, the risks of infection and organ rejection mean the patient's fight for recovery starts immediately; even anti-rejection medication can lead to serious adverse effects. Eurofins' work in immunity is helping to change patient recovery outcomes.

Adverse reactions that can hamper patient recovery are managed by understanding the exact state of a patient's **immune cell function** and using that knowledge to balance the immune system. Experts at Eurofins Viracor developed ImmuKnow® assay, a test that provides a biomarker to help doctors to evaluate a patient's existing immunity levels and to optimise treatment.

In 2015, a study published in *Transplantation* showed that Viracor's assay improves outcomes in solid organ transplant patients. The study showed that not only was ImmuKnow® able to detect the level of immunity provided by cells in immunosuppressed patients, but it ultimately improved patient **survival rates** by helping to optimise immunosuppression. This benefit is critical when balancing the risk of organ rejection caused by insufficient suppression and the risk of infection caused by excessive suppression.

The study also showed that using ImmuKnow® in solid organ transplant

patients helped decrease bacterial and fungal infections, particularly in the critical two-week post-surgery period. It also reduced immunosuppressant drug use and improved patient survival by 13% in the first year post-transplant. ImmuKnow® is cleared for use by the US Food and Drug Administration.

Eurofins Viracor has developed additional, innovative tests with **fast, accurate results** to help surgeons care for their transplant patients. It now works with more than 60% of US transplant programmes and has earned its leading position in the field.

Eurofins is deeply committed to contributing positively to global health. With technological and medical advances leading to continual increases in the number of transplant operations occurring worldwide, Eurofins companies are focused on the development of further specialised and fast-turnaround tests that will play an important role in patient outcomes.



the science behind

ImmuKnow® is an immune cell function assay that detects cell-mediated immunity in an immunosuppressed population by measuring the concentration of adenosine triphosphate (ATP) synthesis from CD4 cells following stimulation. The technology combines cell stimulation, cell selection, and quantification of metabolic markers (ATP) to measure cell-mediated immunity. Since the CD4 lymphocytes orchestrate cell-mediated immunity responses through immunoregulatory signalling, the measurement of CD4 activation reflects the degree of immune function.

For references, please see bibliography

Disclaimer: ImmuKnow® is FDA cleared for the following Intended Use: Detection of cell-mediated immune response in populations undergoing immunosuppressive therapy for organ transplant. Product Insert 2015. Eurofins Viracor, LLC, Lee's Summit, MO.



SALMON UNDER THREAT

DETECTING A NEW CONTAMINANT OF CONCERN

Developing the first test for 6PPD-quinone



Each spring and autumn, millions of salmon migrate along the American West Coast from the Pacific Ocean, returning to their native rivers and streams to spawn. Unfortunately, a significant number of these salmon mysteriously die before they reach their destination, threatening the future of the wild salmon population. Scientists believe that 6PPD-quinone, a toxic compound that is polluting waterways, is to blame. Eurofins Environment North America was the first in the world to develop a commercial testing method for 6PPD-quinone in aqueous and solid matrices.

This problem of wild salmon dying suddenly while migrating has worsened over time and the underlying cause has puzzled experts for decades. However, a late-2020 publication by the University of Washington and other organisations in Science made a major breakthrough in understanding the threats faced by wild coho salmon (*Oncorhynchus kisutch*).

The study has identified a compound that is toxic to the salmon, even at low concentrations, called 6PPD-quinone. This compound begins as 6PPD, a member of a group of compounds known as anti-degradants, which are used as additives to protect tyres and other rubber products from damage from ozone. In doing so, 6PPD is oxidised and transformed into 6PPD-quinone. As the tyres then wear, the worn rubber is continuously deposited on roadways and makes its way into waterways through run-off from rain.

Several follow-up studies have confirmed toxic effects of 6PPD-

quinone to aquatic life, but not all species are affected equally, with some creatures exhibiting a 6PPD-quinone tolerance over 100x greater than that of coho salmon. However, acute toxicity has also been observed in rainbow trout (*Oncorhynchus mykiss*); while coho salmon are isolated to the Pacific Northwest, rainbow trout are found in over 50 countries worldwide. This discovery makes it clear that 6PPD-quinone is a global issue.

As trout is closely related to char, another widespread and economically important genus of fish, there is a need for additional toxicological and transport studies on every continent.

Eurofins Environment North America's world-first commercial method tests for 6PPD-quinone in aqueous and solid matrices, including both sediments and consumer/construction products. Recently, the company pushed the aqueous detection limit from 10ppt (parts per trillion or ng/L) to 2ppt in aqueous samples. This is achieved from only 250mL of aqueous matrices.

the science behind

The technique used exhaustively extracts the contaminant and utilises solid phase extraction (SPE) to concentrate the compound and remove bulk interferences. Analysis is carried out on an LC-MS/MS (Liquid Chromatography-Tandem Mass Spectrometer), utilising isotope dilution quantitation. This approach employs a stable isotope of 6PPD-quinone, which is added to samples prior to extraction. Isotope dilution is considered the gold-standard approach for trace-level environmental analysis because it can loss-correct sample concentrations and mitigate both enhancement and suppression caused by the matrix.

For references, please see bibliography

BUSTING

THE IDENTICAL TWIN MYTH

Milestone research proves genetic differences

Amid abandoned legal cases, high-profile acquittals of serious crimes and paternity suit dismissals, scientists were increasingly questioning the dogma that identical twins could not be genetically differentiated. Only theoretical approaches existed to solve this conundrum, until a breakthrough by a Eurofins company provided an answer to the puzzle of how to tell identical twins apart.

Monozygotic twins derive from one zygote (the fertilised egg) that splits into two, meaning that the basis of the two twins, at first, comprises exactly the same genetic material. By pushing the limits of genetics and forensics, a multi-disciplinary team at the flagship Eurofins Genomics laboratory in Ebersberg, Germany, developed a **world-first methodology** in 2014 that showed what happens next.

Using a whole genome sequencing approach, a Eurofins company demonstrated that rare mutations occur early in the embryonic stage whenever a cell divides as part of the development process. A Eurofins company also provided evidence that, because these changes happen so early on, the mutations can be found in all the DNA of the person, including the DNA of sperm (in the case of the male) or eggs (in the case of a female).

Eurofins' first test case used this new understanding of embryonic genetic mutations to assess identical twin

adult males as the potential father of a child. While one was found to carry an identical mutation to the child, the other was not. These results were used to prove **conclusive paternity**.

The first application of the test in a criminal case supported the Boston Police Department and Suffolk County District Attorney's Office in a serial rape case. The authorities were confident of the guilt of two male offenders; however, one had an identical twin and could not be convicted. Eurofins received saliva samples from both twins, and a crime scene sample. The test delivered a successful result in just two weeks, successfully and unambiguously matching one of the twins to the crime.

Through the use of state-of-the-art chemistry and ongoing process refinements, Eurofins has already cut the test costs by 50%, ensuring the **accessibility** of one of the most important tools in the search for the truth.

the science behind

The process involves ultra-deep Next Generation Sequencing. Researchers first read and decipher the genetic information found in the DNA, before analysing the data collected through intelligent information technology. The focus is on the appearance of specific mutations, also referred to as single nucleotide polymorphisms, or SNPs.

For references, please see bibliography





NEW GENERATION OF NON-INVASIVE PRENATAL TESTING METHODS

Breakthrough genetic disorder determination

For over 30 years, Eurofins laboratories have held leading positions in the research and development of **prenatal screening protocols** to help detect birth defects, genetic diseases and other conditions. For example, Eurofins laboratories were among the world's first clinical laboratories to provide an alpha-fetoprotein screening programme. This test is known for its effectiveness in identifying a risk in the foetus for open neural tube defects.

In December 2016, Eurofins LifeCodexx launched the worldwide first non-invasive prenatal test (NIPT) assay based on a quantitative real-time PCR for prenatal testing of foetal trisomy 21 (qNIPT). The novel and proprietary qNIPT assay allows for significant cost efficiencies and speed over other classical NIPT methods as it is based on polymerase chain reaction (PCR), the most widely used platform within the molecular diagnostic space. This test's significantly lower cost and faster processing time allows more pregnant women to opt for NIPT as a reliable and affordable prenatal testing method.

Similarly, Eurofins Biomnis was the first commercial laboratory in France to offer, in October 2014, an NIPT Next Generation Sequencing test and was also the first to validate the method on twin pregnancies.

In 2017, Eurofins Genoma unveiled a breakthrough in **NIPT** technology, developing, for the first time ever, an NIPT that screens for both de novo-gene mutations, that are not inherited – and inherited single-gene disorders. NIPTs are simple blood tests that were previously only able to screen for aneuploidies, errors in cell divisions which see 'daughter' cells having the wrong number of chromosomes, and microdeletions, which are chromosomal deletions that are too small to be detected by light microscopy and can lead to genetic disorders. More recently, NIPTs also screened for **rare aneuploidies** and segmental chromosome imbalances (gains and losses) in every chromosome in the foetal genome. Not only does this new Eurofins test, GeneSafe™, identify prenatal conditions which traditional prenatal testing may miss, but it also detects over 40 severe genetic disorders

Prenatal tests are conducted during pregnancy to provide information about the health of the mother and her baby. Eurofins companies have pioneered work to develop improved prenatal screening tests for conditions that might affect the foetus, and help inform healthcare decisions before and after the birth.

that might occur in the absence of any family history of the condition.

Eurofins Genoma has significant experience in performing genome-wide cfDNA screening, which can detect rare trisomies. The laboratory is able to distinguish between trisomies that are likely confined to the placenta (resulting in a false positive result which decreases the specificity of the test) from true trisomies (which are confirmed to be in the foetus).

The test is also the first ever NIPT to detect disorders with an increased likelihood associated with **paternal age**, something particularly useful as later-stage parenthood is increasingly common.

Apart from NIPT tests performed during pregnancy, Genoma also developed one of the first **non-invasive preimplantation genetic aneuploidy screening** tests, so a test that is used to exclude chromosomal aberrations of embryos for assisted reproductive technologies.

the science behind

Based on quantitative real-time PCR rather than on Next Generation Sequencing, the qNIPT foetal trisomy 21 test from Eurofins LifeCodexx is not only cheaper, but also significantly faster. Due to different methylation patterns of specific gene regions of the maternal and foetal DNA, positive and negative samples can be distinguished and the test provides rapid results in two to three days under the highest quality standards.

The GeneSafe™ test is intended for patients who had an abnormal ultrasound finding which was suggestive of some form of monogenic disorder. The technology screens for new mutations that are common and cannot be detected by standard carrier screening, as these mutations are not present in the parents.

While traditional NIPT screens for conditions typically associated with advanced maternal age (e.g. Down syndrome), GeneSafe™ also screens for genetic disorders that are associated with advanced paternal age (men that are over 40 years old), ensuring a more comprehensive screen for couples of advanced age.

For references, please see bibliography





Food fraud and the adulteration of what we eat and drink is a global problem and one which is only worsening. Fraudsters are substituting genuine, everyday ingredients with cheaper alternatives that mislead buyers and consumers, and risk allergies, side-effects and even severe harm to human health. The desire to help put an end to such dishonest practices led to the birth of the company that became Eurofins.

Authenticity and origin testing to protect our food and drink

SNIF-NMR®

HOW IT ALL BEGAN



Back in 1981, Professors G. and M. Martin and their teams at the University of Nantes developed a **pioneering process** to detect the chaptalisation (or over-sugaring) of wine, and the enrichment of freshly pressed grapes. In 1987, this SNIF-NMR® method led to the creation of Eurofins. The company has continued to develop the process, now using it to check the authenticity of wines, spirits, fruit juices, flavours, sugar syrups, and vinegar. Together with other tests, the technique makes it possible to verify the geographical origin of products and decide, for example, whether vanillin was chemically-produced or whether it is natural – useful information as the two have a huge price difference.

This technique is still state-of-the-art 35 years after its invention, and is a **gold standard** in its field. SNIF-NMR®

is the official method of analysis for a variety of organisations, including the European Union (EU) and the International Organisation of Vine and Wine (OIV) for wine; the Association of Analytical Chemists (AOAC) for fruit juices, maple syrup, and vanillin; and European Committee for Standardization (CEN) for vinegar.

In addition, Eurofins has committed significant resources to developing a comprehensive reference database for wine, fruit juice, natural flavours and other products. The company continues to pioneer and develop cutting-edge tools to empower customers to fight back against food and beverage fraud.

the science behind

Atoms exist as different isotopes. Sugar in sugar beet, for example, contains different proportions of hydrogen isotopes (1H and 2H). SNIF-NMR® can measure the different hydrogen isotope ratios at each position in a molecule, thereby identifying the addition of sugar from another source. The SNIF-NMR® process allows the quantification, with very fine precision, of isotopic ratios across a molecule. A number of factors affect these, including botanical origin, climate and geography and so the resulting isotopic fingerprint provides in-depth information about the origin of the molecule.

For references, please see bibliography



THE FUTURE OF BEEES

Breakthrough science for supporting better bee health



THE FUTURE OF LIFE



Honey bees travelling from flower to flower are more than just a pretty sight; the species is the world's most important pollinator. Environmental changes including new agricultural practices mean high levels of concern globally about pollinator safety. Eurofins companies are leading efforts to support and protect bee health and reverse the trend. Their innovative approach to homing behaviour studies, for instance, is on its way to becoming part of standard industry protocols.



the science behind

The Eurofins Agrosience Services business line developed an innovative method for fully investigating the **effects of pesticides** on the homing behaviour of honey and bumble bees. Tiny **radio frequency identification transponders** with unique identification numbers are permanently and harmlessly attached to each bee's thorax. The transponders are registered by scanners at the hive entrance whenever a bee enters or leaves, allowing their homing rate to be assessed.

One sample of experienced forager bees is fed different doses of an insecticide and compared to an untreated control sample. The bees are caught and fed within one day. At the end of the study, available data covers the duration of return flights after feeding and the return rate or homing success as a possible

indicator of disorientation. The study design can also be adapted to include realistic field-exposure scenarios and to observe delayed effects for the remaining flowering period of the crop. The Eurofins method provides data crucial to understanding the impact of the insecticide and any direct links between test types and their impact on bee health. The method was also successfully applied in a field study with bumble bees. Furthermore, since 2021 Eurofins Agrosience Services has participated in a novel approach to count bees entering and leaving the hive. The bee counter is developed with a start-up and can not only count bees entering and leaving the hive but also notes if a bee brings pollen into the hive.

Just as innovatively, companies of the network pioneered a field application method to assess the impact of

abraded dust from pesticide-treated seeds on honeybees. Varying particle size makes the dust inherently more difficult to test than liquid substances; however, abraded pesticide particles or contaminated dust contain high concentrations of pesticide and can prove toxic to bees actively collecting pollen and nectar. Finding a solution was imperative, but previous trials proved labour-intensive and complicated and were affected by numerous factors including wind direction. A Eurofins company developed a **purpose-built dust applicator** which improved risk assessment methodology and opened up a range of field trials to investigate potential side effects on bees and other organisms.

Eurofins companies also take an innovative approach to breeding methods for honey bees, using genome-wide DNA analysis. Working alongside

the Institute for Bee Research in Hohen Neuendorf, Germany, and the department of Molecular Biology and Genetic, Aarhus University, Denmark, they have sequenced bee genomes using Next Generation Sequencing to identify genomic loci that regulate a worker honey bee's ability to detect and remove diseased or infested brood, a key 'hygienic behavior' to protect the colony against the Varroa parasite. They have also been able to characterise genetic variants that impact this trait. A new testing solution from Eurofins, launched in late 2023, can help to breed more resistant colonies by selecting bees that exhibit this hygienic behaviour, leading to healthier bee populations that support functional eco-systems.

Eurofins' key role in safeguarding bee health includes solitary, honey and bumble bees. Quality bee breeding is aided by testing queens and drones for fertility, reducing stressors and optimising nutrition. To aid selection of disease-resistant bees, a Eurofins company analysed the whole bee genome using Next Generation Sequencing to identify single nucleotide polymorphisms (SNPs), small natural changes, in the bees' DNA. Bee variants are sorted according to the SNPs, using DNA chips developed to characterise gene variations in bees with known parasite resistance, productivity and docility. This allows beekeepers to identify the genetic basis for these positive traits and avoid time-consuming selection.

For references, please see bibliography



HELPING TO STOP ZIKA IN ITS TRACKS

Rapid real-time Zika test results



The summers of 2015 and 2016 could be remembered as the summers of Zika, with thousands of confirmed and suspected cases of the virus, real fear among travellers to affected regions, and withdrawal from the Rio Olympics by dozens of worried athletes. Eurofins companies responded with the fast-track development and emergency launch of a test to help stop the spread of this heart-breaking and virulent virus.

The Zika virus was first linked to a fast growing number of cases of **microcephaly** (a severe neurodevelopmental disorder resulting in smaller-than-normal head size) in newborns, particularly in Central and South America in 2015. Global concerns grew when travellers to these areas, and to others endemic to the virus, began to test positive for Zika. Spread primarily through *Aedes aegypti* mosquitoes, Zika virus can also be passed on by some forms of sexual contact.

Determined to assist their clients and patients prior to the onset of the mosquito season, Eurofins Biomnis and Eurofins Viracor became amongst the first commercial laboratories to develop and offer real-time polymerase chain reaction (PCR) tests for Zika to hospitals and health care providers. Subsequent outbreaks in French Guiana and the West Indies in late 2015/2016 required large-scale testing. Working with the French National Reference Centre for Arboviruses, Eurofins Biomnis obtained authorisation in late December 2015,

making the test swiftly available for use at national and international levels by January 2016. Eurofins Viracor, in partnership with the US Food and Drug Administration, obtained emergency use authorisation in July 2016 in the USA.

Eurofins Zika assays test human blood serum, blood plasma, urine, saliva, sperm and amniotic fluid from individuals presenting Zika-virus signs and symptoms. It is designed for those who have a history of living in, or travelling to, affected areas. This **rapid-response** test gives doctors and healthcare professionals a result within just eight to twelve hours of receipt of the sample by a Eurofins laboratory.

The assay's exceptional sensitivity safeguards against cross-reactions with other viruses in the same family (Flaviviridae), including Dengue, Japanese encephalitis, West Nile, and St. Louis encephalitis, or with other viruses known to cause similar clinical symptoms, such as Chikungunya.

the science behind

Eurofins Biomnis and Eurofins Viracor's real-time PCR tests for Zika virus are designed to detect and measure ribonucleic acid (RNA). RNA is extracted from the specimen and reverse transcription of viral RNA to DNA is used. With the help of the polymerase chain reaction (PCR), a laboratory method used to make very large numbers of copies of short sections of DNA from a very small sample of genetic material, the number of DNA copies obtained is amplified and enables specific genes of interest to be detected or measured.

For references, please see bibliography

A PIONEER IN DIOXINS DETECTION

Detecting one part out of 1,000 billion is not the limit

To protect human health, understanding the levels of dioxins in the environment and the food chain is imperative. Eurofins is at the forefront of global efforts to limit our exposure to dozens of dangerous chemicals.

the science behind

Generally, POPs are substances containing chlorine, fluorine or bromine within their molecules. One of the most typical examples are PCBs. These are a class of chlorinated aromatic hydrocarbons which consist of 209 individual components. They started being commercially produced in 1929 but were recognised to be harmful in the 1970s and have since been banned or severely restricted. It is estimated that around two million tons of PCBs have been produced, about 10% of which still remain in the environment today. Twelve of the PCBs are similar, in terms of molecular biological behaviour, to dioxins and are known as dioxin-like PCBs.

For references, please see bibliography

Dioxins and the similar polychlorinated biphenyls (PCBs) are persistent organic pollutants (POPs) – toxic chemicals that persist in the environment and accumulate in the food chain. Even at the level of 1/1,000,000,000 to 1/1,000,000 of a gram, they are considered to pose a **serious hazard** to humans and the environment. National and international treaties and directives have been passed to minimise or stop their circulation.

A Eurofins company has led the way in testing and analysis to aid understanding of the levels of POPs to which we are exposed. In 1985, the company was one of the first to start dioxin analysis and, by 2010, became the very first laboratory in Europe

able to test for all the **POPs** identified in the 2001 Stockholm Convention including full analysis of all 209 PCBs with maximum possible separations within one method. This is of particular importance for the fish oil industry, as PCBs and dioxins are lipophilic (soluble in fat, not water), therefore accumulating in fat and oil over the food chain.

Dioxin analysis is important across a variety of different matrices, including food and feed, human blood and tissues, animals, plants, air, industrial products, residues, sludge, and water. Eurofins companies analyse all of these. However, dioxin analysis is extremely difficult and the requirement for the detection of very low

substance levels (one part out of 1,000 billion others) and the distinction of each substance among the very similar dioxin and PCB molecules necessitates the use of highly advanced equipment. Eurofins laboratories are amongst the world leaders regarding full process automation, method development and advances in gas chromatography and mass spectrometry in order to make this type of analysis reliable and helpful for hazard control as well as scientific research. Extraordinary investment in research and development has helped Eurofins achieve a market leading position globally, and the laboratories now analyse over 100,000 dioxin samples per year, performing the highest throughputs worldwide. Eurofins has

released almost 400 publications on dioxins and POPs testing to date.

One of the sources of dioxins can be waste incinerators. These burn substances containing chlorine at high temperatures, creating the conditions for dioxin formation. For this reason, emission gases are sampled and analysed to ensure that industrial emissions are as low as required and the respective gas cleaning equipment is functioning properly. Eurofins companies are world leaders in the area of sampling and analysis of emission gases, including analysis of the equally toxic brominated dioxins.

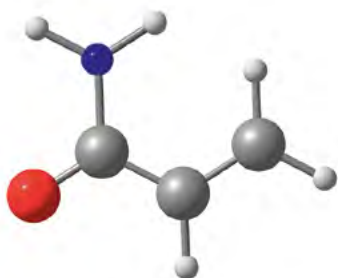




First to detect acrylamide in food

ACRYLAMIDE ON THE TIP OF EVERYONE'S TONGUE





Some rather unusual discoveries in Sweden in the 1990s led to the finding that acrylamide is not only toxic, but also present in a high number of foodstuffs. Acrylamide is understood to be probably carcinogenic to humans and both the World Health Organization and the Food and Agriculture Organization of the United Nations view the levels of acrylamide in foodstuffs as a major concern. Eurofins has played a leading role in detecting and tracing occurrences of the substance and working to increase understanding of its dangers.



Soon after the walls of a new train tunnel through a mountain range in southwest Sweden were sealed in 1997, fish in nearby waterways started to die and cattle showed signs of paralysis. An immediate investigation by the authorities saw the Eurofins team in Lidköping, at that time AnalyCen AB, perform a large number of initial tests on water, soil and sediment. The cause of the problems in fish and cattle was found to be a **significant leak** of un-polymerised acrylamide and methylolacrylamide (acrylamide is the starting material to produce the non-toxic polymer polyacrylamide).



Further investigations showed widespread contamination in creeks and groundwater across the area, and the grouting material used in the tunnel construction was found to contain both acrylamide and methylolacrylamide. Concerns grew for both the local population and for

the construction workers who had been exposed, but tests conducted by Stockholm University's Department of Chemistry showed 'relatively high' background levels of acrylamide even in their control group. This meant there had to be another **exposure source**.

This source became clear during 1998, when the Eurofins team in Lidköping completed comprehensive feed and food analysis, making the **breakthrough discovery** that acrylamide was forming in heated food. A partnership with Stockholm University saw a huge project involving extensive analysis and the frying, baking and boiling of all kinds of foodstuffs to better understand what was happening. The team discovered that when the amino acid asparagine – found in many vegetables, and in higher concentrations in some varieties of potatoes – is heated to high temperatures in the presence of

certain sugars, acrylamide can form. High-temperature cooking methods such as frying, baking or roasting were more likely to produce acrylamide, but only when heated to a temperature above 120 degrees Celsius (248 degrees Fahrenheit).

The team continued to work over several years to further identify the causes of acrylamide formation, and to refine the analytical methodology and advise food producers and consumers alike. Across Eurofins globally, there are now more than 25 laboratories able to support the food processing industry with this analysis. With acrylamide formation still a topic of concern, Eurofins teams continue to innovate and refine practices in this field, such as by incorporating fully automated sample preparation robots into the process.

the science behind

Acrylamide, a colourless, non-volatile crystalline solid, is a raw material (monomer) in the production of polyacrylamide polymer, used as a coagulant in wastewater treatment and to clarify drinking water, as well as a grouting agent for the construction of dam foundations and tunnels. Potential contamination from the environment meant that reducing the levels in food became even more critical and the European Commission for Food Safety has adopted a recommendation that acrylamide levels in food are monitored with a view to reducing them.

For references, please see bibliography



TESTING FOR TRACEABILITY OF MEAT

Clarity and confidence for the food chain

FROM FARM TO FORK DOWN TO INDIVIDUAL ANIMALS

Food safety crises, from bovine spongiform encephalopathy (BSE, or “mad cow disease”), to the European horsemeat scandal, have rocked the food industry and driven consumers to demand safer food products and reliable information about sources and production processes. Eurofins companies' innovative testing approaches have helped provide true “farm-to-fork” traceability and much-needed confidence to the sector.

BSE was first identified in cattle in 1986 and, unfortunately, necessitated the slaughter of over four million animals in the UK alone. In 1996, the disease was acknowledged to have jumped the species barrier to humans, via the consumption of infected beef and beef products. The first cases of a new variant of Creutzfeldt-Jakob disease (CJD)

slaughter. Large-scale adoptions of the technology for DNA traceability projects included the Pays de la Loire region of Western France.

Technologies have continued to evolve, and the Eurofins network is still leading the way in rapidly reacting in times of crisis. For example, a DNA

were registered and over 220 people are believed to have succumbed to the disease in Western Europe. In addition to these human and animal fatalities, the global news headlines resulting from the BSE scandal and other food safety crises cause pressure across the supply chain and highlight the very real importance of being able to reliably trace the origin of meat products.

The technology to achieve this lagged behind until, in 2001, Eurofins developed its innovative patented method, **Eurofins TAG**. For the very first time, genetic fingerprints were used for the precise identification of cattle in the food industry that had been tested free of BSE, with each individual animal matched to one genetic profile. Simple to administer, reference samples were taken from animals before they reached the slaughterhouse, and the information was stored in a database and compared with control samples taken post-

chip-based method was validated in 2015, in the wake of the horsemeat crisis in 2013. Consumers, retailers and producers were demanding the verification of product composition, and proof of its compliance with relevant regulations and that it was entirely consistent with the claims on the label. The DNA chip allowed for the identification of animal species in food products, including pig, chicken, rabbit, cat, sheep, and goat, as well as horse. In 2024, the laboratory Eurofins Biologie Moléculaire France, based in Nantes, replaced the DNA chip with a set of newly developed multiplex Real-time PCR tests for the detection of up to 12 animal species in meat products, compound food samples, and animal feed. Contrary to the DNA chip-based method, the Real-time PCR technology includes an internal inhibition control, meeting the most stringent requirements in terms of quality assurance.



the science behind

Eurofins TAG is based on the multiplex analysis of a set of 11 microsatellite markers recommended by ISAG (International Society for Animal Genetics). A unique genetic fingerprint is obtained for each animal and can be used for traceability studies of BSE-tested animals of known origin or paternity testing.



For references, please see bibliography

THE FOREFATHERS OF PESTICIDE TESTING

First multi-pesticide detection method in Europe

Pesticides were launched in the 1930s, and the protection they afforded crops was lauded. However, the hundreds of pesticides sold worldwide, with more developed each year, plus cases of the use of banned pesticides like DDT and other persistent organochlorine compounds, mean ongoing pressure to ensure that foodstuffs are safe for consumption.

It was only in the 1960s that concerns for the impact of pesticides on human health began to be raised, but the first methods devised for pesticide analysis were slow, expensive and imprecise. This all changed when Dr Wolfgang Specht recognised that all of the pesticides in use were relatively similar, and that analysis of several of them at the same time would be significantly more efficient as well as more effective. In 1964, he launched the ground-breaking S19 pesticide analysis test as the first ever multi-method in this area.

Used to analyse the contamination in foodstuffs caused by insecticides and other toxic substances, the multi-residue S19 method later evolved to become the official European enforcement method, and the laboratory that developed it – part of the Eurofins family since 2000 – is the recognised **Competence Centre** in the field of pesticide analysis.

Demand for multi-residue methods in plant protection analysis is huge, but more than half of existing pesticides can be assessed with just one method.

Moreover, multi-residue methodology means not only detection, but also direct simultaneous quantification, of residues in order to prove that maximum residue levels are not exceeded. This makes significant contributions to consumer safety.

In 1979 the first method using S19 was published – for tobacco. This test's **real innovation** was the first-ever use of gel permeation chromatography (GPC), allowing the separation of the pesticide analytes from the matrix, in particular fat, by size exclusion via a soft gel. At

first, around 200 molecules, mainly organochlorine and organophosphorus pesticides which were primarily used during that time, could be analysed; Eurofins is now able to analyse many more pesticides and their metabolites with modern methods, processing over half a million samples each year.

Thanks to Eurofins' pioneering spirit, a portfolio of validated methods for the analysis of residues in fruit and vegetables, baby food, tea, spices, herbs, tobacco and animal originated matrices is now available.



the science behind

The GPC technique is the most notable step within the Specht-Method and this was further evolved and utilised for the clean-up of polycyclic aromatic hydrocarbons, ubiquitous environmental pollutants generated mainly through the incomplete combustion of organic materials.

Their testing requires the removal of fat without removing fat soluble pesticides. To do this with a solvent is not possible, however GPC works instead on assessment of the similar molecular size of pesticides.

Further developments have seen a new extraction and partition step introduced into the well-established multi-residue DFG method S19, replacing dichloromethane with ethyl acetate/cyclohexane. In addition, working processes were further refined. The latest development is the reduction in use of solvents in order to protect the environment and save costs.

For references, please see bibliography



35 YEARS OF DATA FOR LIFE

Building rich databases before Big Data



For more than three decades, Eurofins has been considered a leading provider of analytical services globally. The Group performs hundreds of millions of tests each year to establish the safety, identity, composition, authenticity, origin, traceability, and purity of biological substances and products. Its extensive databanks offer years' worth of information about pharmaceuticals and food and their properties.

Many of the tests that Eurofins performs around the world rely on one of the company's extensive **proprietary databases**, not simply to compare results to, but very often to obtain the results themselves.

Eurofins' **DNA fingerprint** database, for example, contains unique identifying characteristics (the "fingerprints") of foodstuffs and enables proof of authenticity that had not previously been available. Not only does this prevent unscrupulous suppliers gaining an unfair market advantage, but it also prevents them endangering the health of the consumer.

Eurofins developed its proprietary DNA fingerprint databases for several specific analytical requests (e.g. **basmati / fragrant rice authenticity** or determination of different pine nut species), using proprietary or published DNA fingerprint methods and reference samples which were made available by authorities or via Eurofins' laboratory network and its customers.

The databases allow identification of pure and mixed samples, but also of the presence of as-yet unknown or unapproved species or varieties. The testing methods meet the need for traceability – a major requirement of EU food legislation – and can be adapted very fast to the needs of the market, for example if new species or varieties are approved by authorities and reference material is available.

Another significant breakthrough in proprietary databases came with the launch of Eurofins' BioPrint™ databases in 1999 to improve and optimise the choice of drug candidates. Much more recent, building on that original innovation and to incorporate internally developed artificial intelligence (AI) tools, Eurofins DiscoveryAI SAFIRE™ (Suite of ADMET Predictions for In Silico Refinement and Evaluation) was launched in 2024 – an advanced platform that leverages the BioPrint dataset, artificial intelligence and machine learning to accelerate drug discovery. SAFIRE utilises data from Eurofins Discovery's



the science behind

Information about the origin of a food product is often encrypted in its chemical composition, and rapid developments in science and technology over the last few decades allow its analysis and interpretation. Two of the most important techniques within DNA fingerprinting are DNA fragment length analysis and microsatellite or short tandem repeat (STR) analysis. DNA fragment length analysis considers changes in the length of a specific DNA sequence to indicate the presence or absence of a genetic marker or the presence of a specific variant of a genetic marker. Eurofins used this technique to successfully detect a species of poor-tasting pine nuts which had triggered 39 biotoxin notifications in the EU Rapid Alert System for food and feed. STR analysis is used to compare specific areas on DNA from two or more samples. This technique, again based on Eurofins' extensive proprietary database of specific DNA fingerprints, was used by the company to prove the authenticity of Basmati rice when the market was flooded with cheap imitations.

The BioPrint™ project was started with the hypothesis that the in vitro pharmacological profiles of new drug candidates generated in Eurofins' laboratory could act as a fingerprint, capturing information on the in vivo activity of the compound. The company found that hierarchical clustering of the drug and reference compounds based on their in vitro pharmacological profiles, achieved the grouping of many by their therapeutic areas or biological actions; for example, antidepressants clustered with other antidepressants, and antifungals with other antifungal drugs. Using this "fingerprint", choices about a drug candidate's potential therapeutic use, and adverse reactions in the context of all the drug and reference compounds present in the database by performing simple profile similarity analysis to identify "neighbour" compounds.

The backbone of the QTA® solution is the comprehensive approach taken in model and application development, technical consultation, and full service support. QTA's unique data treatment methodologies are being applied to calibration models for spectroscopic qualitative analysis applications. The proprietary database and algorithms are dynamically maintained for superior accuracy and precision, with primary data generated using industrial standard methods and stored in a highly-secure central server.

For references, please see bibliography

BioPrint database, curated over years of meticulous research and data accumulation. The BioPrint database comprises a large and homogeneous set of experimental data, generated in-house and containing more than 2,400 compounds, including marketed drugs, compounds which failed in clinical trials, and reference compounds. Each compound has hundreds of pieces of information stored, with the BioPrint™ database covering in vitro assays as well as in vivo characteristics such as drug reactions, pharmacokinetics and therapeutic indications.

Each compound has hundreds of pieces of information stored, with the BioPrint™ database covering in vitro assays as well as in vivo characteristics such as drug reactions, pharmacokinetics and therapeutic indications. On average there are 400 records for each compound, meaning more than 1 million records are stored. High quality and extensive datasets, combined with modelling and mining tools, place new drug candidates in the context of well-understood drugs.

This allows scientists to **anticipate adverse drug reactions** and supports lead compound characterisation and prioritisation (a lead is a possible drug candidate but which may still have suboptimal structure and characteristics).

A further development is a cloud-based proprietary database from Eurofins QTA that provides a major advancement in the way customers perform their infrared (IR) analysis.

IR analysis is a method to measure at what wavelength and how much light is absorbed by a material type and at what intensity light passes through it. Individual wavelengths interact with different compounds in different ways. This allows chemists to use the data to map the physical and chemical properties with quantitative and/or qualitative data.

The QTA® methods provide a simple interface for non-skilled users, thereby overcoming the challenges of traditional infrared analyses which have to be developed and maintained

by experienced spectroscopists or chemometricians. QTA provides clients with a uniform analysis method that can be utilised by any non-technical personnel across multiple locations, reducing the human error that can occur when analysing with traditional chemistry methodology.

Instruments at the testing site scan a sample and send the light spectra of the sample to the secure server via the internet. The analysis and data interpretation are completed within minutes on the server, and the results are returned in real time. Common sample matrices include food products, chemical manufacturing, oils, agriculture, aquaculture, and dairy. Eurofins QTA pushes the boundaries of where NIR is able to be used in order to introduce this analytical solution into new applications and emerging industries using benchtop (i.e. in the laboratory) or in-line (i.e. on the production line) systems.



FINDING ADULTERATION

Leading the way with non-targeted screening

EVEN WHEN NOT LOOKING FOR IT

When you search for something specific, that's probably all you are likely to find. By removing the constraints of targeted analysis and searching without restriction, Eurofins has opened up a whole world of new possibilities.



Eurofins' evolution of non-targeted screening methods is one of the company's most exciting developments. Teams from across the company have pioneered wider screening techniques covering a range of sectors, from food and pharmaceutical, to environment and water. A very prominent example of an unexpected, and hence missed issue that evolved into a scandal was the addition of the industrial chemical melamine found in baby milk in China in 2008. Another was the addition of horse meat to beef in Europe in 2013. Such fraudulent practices were never expected and so were not explicitly searched for. Non-targeted screening tests are enabling a whole variety of industries to better guard against health crises.

High-resolution nuclear magnetic resonance (NMR) testing is one example for such an **untargeted screening approach**. Eurofins' method is now able to provide additional insights to those offered through conventional approaches. This

investigative technique allows for the detection of hidden anomalies, such as the addition of sugar to honey, or the mislabelling of geographical or botanical origins in the composition of a complex matrix. The approach can be applied to liquids or extracts from solid foods, requiring only limited preparation of the sample.

Another pioneering technique from Eurofins' laboratories is a new non-targeted screening method, based on **16S rDNA screening** and Next Generation Sequencing. Used to profile the microbiome of food, pharmaceutical and environmental samples, the test focuses on identifying the micro-organism communities present, helping to recognise a broad spectrum of bacteria. For example, the test can be used for the control of fermentation processes or the analysis of biofilms (a thin microbial layer). The tests require no assumption to be made about a given species of interest, and all species included in the database can be identified in one test.





the science behind

These sophisticated testing methods also have numerous applications in the environmental world. Eurofins Miljø has, in cooperation with University of Copenhagen, Denmark, developed the two untargeted methods, ChemFingSOIL® and ChemFingAIR™, to address environmental hydrocarbon pollution in soil and indoor air.

The first commercial project to use ChemFingSOIL® was launched in February 2017. In 2021, it was followed by ChemFingAIR™, which uses advanced data modelling to determine the contribution and flux of volatile pollutants from underlying soil pollution to the indoor environment of a house. Similarly, Eurofins Miljø also participates in several public funded R&D projects with the aim of leveraging non-targeted chemical fingerprinting and screening through HRMS (high resolution mass spectrometry) to industrial application in the environmental industry. The company is currently taking part in the AQUAPLEXUS project, which aims to develop and standardise testing

methods to ensure drinking water is safe for human consumption.

In 2020, Eurofins Analytics France launched the ongoing TOFoo® (True Organic Food) project, a collaborative research project supported by the French State, which aims to develop analyses to ensure the integrity of organic food products. Through the project, the Eurofins company and its research partners have developed a multi-technical and non-targeted screening method to distinguish organic from conventional UHT milk and tomatoes. The method relies on an extensive database of organic and conventional reference samples, collected over several years as part of the project, which was used to build statistical models based on artificial intelligence. Thanks to the non-targeted nature of the technique, it is able to reveal intrinsic characteristics linked to specific farming practices in the organic sector, without narrowing analyses to a particular substance or family of molecules.

And this is only the beginning. Eurofins Analytics France is conducting further research on different organic food products, with other analytical techniques currently underway, while other Eurofins laboratories globally are also developing a whole range of other non-targeted screening techniques. Eurofins' insights are helping to ensure that consumers and the wider world at large are fully informed about exactly what is in the products we all consume.

Eurofins' techniques are based on multivariate data analysis, multiple analytical approaches and extensive use of computer models.

NMR is carried out on the protons or carbon atoms of the molecules to be analysed, a full resonance spectrum becoming available within minutes. This technique allows the non-targeted detection of all organic compounds, and the NMR profile can be regarded as a unique fingerprint of the sample. Databases of authentic samples allow the establishment of statistical models for checking the integrity of foodstuffs.

Eurofins' launch of its Next Generation Sequencing test service has opened up non-targeted bacterial screening for food and feed testing. Used for raw or processed food matrices, it provides a view of flora in the product thanks to bioinformatics and database tools. 16S rDNA is the commonly used and recommended DNA target region for bacterial species identification.

ChemFingSOIL® and ChemFingAIR™ both rely on extensive databases of soil and air pollutants, combined with chromatographic data, advanced statistical analysis, and modelling of GC-FID and GC-MS files (Gas Chromatography with Flame Ionization Detection and with Mass Spectrometry respectively). While ChemFingSOIL® is built on the chemical pattern of hydrocarbons in soil, ChemFingAIR™ compares the fingerprint of volatile compounds in soil air with fingerprints of indoor air.

For references, please see bibliography



MODERN FARMING IS PRECISION FARMING

Farming and herd improvement testing – fast and efficient



Dairy farming has been undergoing evolution since the 1960s. Large and highly-specialised farms are now the standard, and dairy farmers are ever-more focused on herd improvements and introducing value-added tests to their farm management. Through its highly- automated milk laboratories, Eurofins is helping farmers produce increased and higher-quality yields from healthier herds.

Eurofins has expanded its services from standalone testing to supporting the whole value chain at the farm, in order to help dairy farmers to make informed decisions in favour of food safety and animal welfare. The company has developed a range of tests to provide insights into the origin of milk, the total yield, cow's health, feed efficiency, environmental impact, quality, and adherence to all relevant regulations. Eurofins' approaches have reduced test times from weeks to just days and hours, providing results for more than 20 different parameters for the management of the farm.

The dramatic **milk yield increases** achieved since the 1960s are

testament to the effectiveness of this precision farming and precision testing, with the average yield increasing from 4,200 litres per year to over 10,000 litres per year for an average dairy cow in Denmark.

Testing itself has also developed markedly, with initial tests providing only limited, simple analysis of fat content with moderate accuracy. Eurofins now offers multiple testing, at **high throughput** in automated specialised laboratories and delivers approximately 200,000 results each day from 30,000 milk samples. Tests can be conducted on an individual cow, or on the milk bulk tank, and routine tests now include fat and

protein to evidence quality and assess price. The **somatic cell count** – consisting primarily of the white cells – are measured too, as increases in these can be a sign of a response to pathogenic bacteria and lead to illness or disease in the animal.

Eurofins is also able to run bovine diagnostics (ELISA antibody testing) to measure factors such as beta-hydroxybutyrate, which indicates ketosis or possible energy deficiency of the cow, and Paratuberculosis (Johne's disease), which is a contagious and chronic disease reducing the milk yield. Other factors covered in the diagnostics include Bovine Viral Diarrhoea, which is a

highly contagious infection causing diarrhoea; Infectious Bovine Rhinotracheitis, which causes generalised systemic infections, and Bovine Spongiform Encephalopathy – commonly known as BSE – a fatal central nervous system disease. These all have significant implications for the animals' welfare, fertility and yield, as well as creating potential issues around food safety and adherence to legislation.

Eurofins is already working on more dynamic and interactive testing to include differential somatic cell count, genetics, environmental information, feed efficiency, and differentiation of spores.

the science behind

Four primary tests are used to test milk:

(i) the Fourier Transform infrared test (FTIR), using mid infrared spectrum to predict more than 18 different parameters, including fat, protein, lactose, urea, casein, ketosis, freezing point depression, solids, adulterants;

(ii) flow cytometry for detection of somatic cells in milk. Flow cytometry is a technique used for counting and characterisation of particles and cells, and can be described as "the measurement of cells while they are flowing";

(iii) flow cytometry for detection of the total count of bacteria in milk. Bacteria are counted through specific staining of the DNA and the use of a fluorescent marker;

(iv) diffusion testing for screening for inhibitors in milk. This test looks for antibiotic residues in milk and is performed using 96-Well Microtiter™ plates with agar and a coloured pH indicator. Spores from the test organism *Bacillus stearothermophilus* var. *calidolactis* will have a reduced growth when an antibiotic is present in milk. Results are qualitative and either positive or negative.

For references, please see bibliography

BEATING DRUG-ASSISTED RAPE—

Identifying knock-out drugs in hair

FORENSICALLY

Detecting drugs in hair has been possible since the early 1980s using forensic toxicology, a useful diagnostic tool in proving past drug use or exposure. Two Eurofins forensic toxicology laboratories – Toxlab in Paris and ChemTox in Strasbourg – were involved in the ongoing evolution of testing techniques for several years and together achieved an important breakthrough that assists the judicial system and victims of drug-assisted crimes.

Psychoactive substances are being used to facilitate crimes such as theft and sexual assault. Administered surreptitiously and without the knowledge of the victim, such crimes often involve drugs with a very short half-life. The drugs, including **hypnotics, sedatives and antihistamines**, can rapidly impair consciousness and cause loss of memory of events that occurred after



drug intake. This meant laboratories and law enforcement agencies were struggling to assess drugs with a short 'life', and victims of crimes such as rape, sexual assault or theft were often left not immediately or fully understanding what had happened to them. Delays of more than two or three days in reporting such crimes only compound the problem, as most of the drug molecules were undetectable through the techniques classically used in forensic toxicology (gas and liquid chromatography). Presenting a solution, Eurofins used **liquid chromatography** coupled with **tandem mass spectrometry** and with triple-quadrupole, for the determination of these molecules in biological fluids and, more particularly, in hair. Essentially, they were able

to scientifically and conclusively determine the presence of a drug in hair samples provided as evidence in drug-facilitated crimes, even when there was a delay in the submission of evidence for analysis. It could also show that the drug detected was only ingested once on the date of the crime, eliminating defences that the victim was a regular user of that drug. Given the variety of analytes, the

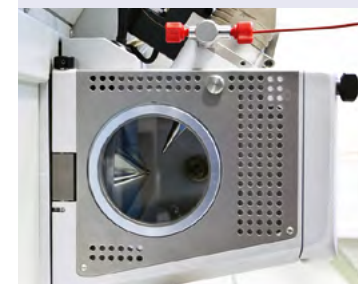


limited amount of hair samples, and the need for a fast result, the company limited the number and the complexity of the extractions, and focused on the technique of **selected reaction monitoring** (see right). Once the technique had been validated, Eurofins helped the French Ministry of Health and the French Ministry of Justice to publish a circular outlining how to obtain biological samples (blood, urine and hair) in such cases. They also participated actively in the organisation of external quality controls and the creation of a committee on drug-facilitated crimes, and they continue to provide their data to the French Health Agency to help them build an accurate epidemiological database.

the science behind

Traditional radio-immunoassay testing evolved into liquid chromatography testing coupled with tandem mass spectrometry to meet the requirements of the Courts of Justice for analysis by two independent methods. Liquid chromatography coupled with mass spectrometry is applied in forensics mainly for the analysis of thermally-unstable molecules in complex samples such as biological fluids. A triple-quadrupole mass spectrometer is a tandem mass spectrometry method in which the first and third quadrupoles act as mass filters, and the second causes fragmentation of the analyte through interaction with a collision gas. Selected reaction monitoring is a method used in tandem mass spectrometry where an ion of a particular mass is selected in the first stage and an ion product of a fragmentation reaction of the precursor ion is selected in the second stage for detection. Having information on the mass of the molecule (ion) itself and its fragmentation products allows for unambiguous identification of molecules.

For references, please see bibliography





HIGH-FLYING INNOVATION

Unconventional inspiration for DNA synthesis

Ready access at realistic cost to short DNA strands is crucial to many areas of scientific investigation. Turning to aerospace for inspiration, Eurofins made a vital ingredient more affordable and, in doing so, helped enable huge numbers of scientists elsewhere continue with valuable research.



the science behind



Nucleotides are the building blocks of life, and a form of these – oligonucleotides (or oligos) – are short, synthesised DNA strands required by scientists for almost all polymerase chain reaction (PCR)-based research. PCR sees stretches of synthesised DNA (oligo synthesis) amplified by creating a number of identical or near-identical copies and is used in a huge set of basic and applied research applications, across all areas of life science and a whole range of genomic investigations. In most cases, the synthesised oligos are used to determine whether a searched-for DNA sequence is present in a product or biological system.

The uses for oligos are broad but, when Eurofins was founded in the 1980s, oligos' high price rendered these **critical research materials** unobtainable for all but the most well-funded scientists. Eurofins Genomics US developed an innovative new platform enabling the production of oligos at an attractive price-point,

essentially democratising DNA and making oligos available to everyone for research.

Since that time, there have been many different iterations and advancements in **synthesis technology**, culminating in a further developmental leap in 2017. A major shortcoming of oligo synthesisers was their wear and the frequency of breakdowns with associated downtime. Here, Eurofins Genomics US turned to an unlikely field for the model – aerospace engineering – completely rethinking the DNA manufacturing process and, once again, succeeding in reshaping the marketplace.

Building a **DNA synthesiser** to manufacture these oligos is almost like building a small jet engine – both utilise the science of air pressure, air flow, vibration, fluidics, and thermodynamics, inside complex, mechanical engineering. But it was the adoption of aerospace engineering's

reaction to downtime – the sector considers it unthinkable – that facilitated the greatest step-change.

To engineer its next generation DNA synthesiser, Eurofins Genomics US partnered with a commercial-grade, aerospace engineering company. The result was the production of high-quality oligos in a shorter time. Also, owing to the incredible power and speed of the machine, the Eurofins team raised the ceiling on scalability and volume, its **innovative design** making the production of even extremely small volumes of DNA affordable.

Not only has this new device filled a critical gap in today's synthesis environment, but the advancement has the potential to help fuel deeper investigations in the biological sphere, revealing important new insights and ushering in a new era of research and scientific progress.

Scientists who buy oligonucleotides demand several conditions: high quality and reliability, rapid turnaround time, low cost, and the right quantity. Unfortunately, previous DNA synthesis technologies failed to accomplish all of these at once. Traditional high-throughput oligonucleotide synthesisers use a lot of reagents and make more DNA than is required in most applications. By contrast, emerging microarray or chip based methods generate too little DNA that, besides requiring additional downstream work before being useful, is only suitable for specific applications, predominantly, gene synthesis.

For references, please see bibliography

TESTING FOR TAILORED CANCER TREATMENT

Empowering cancer research through innovation

Cancer is a complex and multi-dimensional disease, driven by genetic instability and influenced by a multitude of factors. Each patient is unique, and each tumour ever-changing, yet treatment historically has taken a generalised approach with limited success. Advances by Eurofins are helping researchers to “personalise” cancer treatment and better target care for better outcomes.

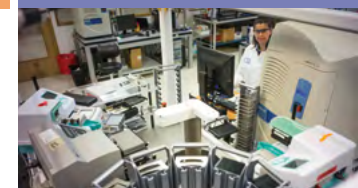
The Eurofins Pharma Discovery Services team in the US utilises its OncoPanel™ Cell-Based Profiling Service comprising more than 300 genomically-diverse human cancer cell lines across 18 different tissue types. The service screens and profiles **anti-cancer compounds** in drug discovery, and also predicts potentially beneficial responses.

This means that drug candidates can be assessed in the very early stages of their development. By statistically correlating the genetic profile of each tumour cell with its response to the potential drug, or drugs in combination, predictive biomarkers can be identified. Drug discovery researchers can now utilise this service to determine which genomic features may predispose patients to successful treatments or identify **potential resistance mechanisms** that may diminish the effect of treatment.

Additionally, Eurofins Discovery offers the OncoPanel™ service using 3D cell culture techniques that add even greater insights by more closely modelling tumour structure. The cells are grown as three-dimensional tumour spheroids and designed to mirror key attributes of human tumours. Innovative, high-content imaging allows the comparison of 2D and 3D cell cultures to assess the efficiency and effectiveness of various drugs in penetrating each tumour.

Truly personalised insight into drug discovery means benefits for patients through the possibility of **faster and better results** from cancer treatment plans and the potential for health care services to reduce waste through inappropriate or inefficient drugs and therapies.

the science behind



High-content imaging enables simultaneous detection of a potential drug's ability to inhibit tumour cell growth and other biological responses. Depending on data requirements, the services can be performed as short and long-term cell culture, as well as 3D models. Such 3D cultures, as opposed to standard 2D models, are particularly interesting, because they take account of the fact that a tumour in real life is not a flat assembly of cells. The 3D cultures hence mimic morphological and phenotypic characteristics of real tumours. They are grown in a way to ensure that the majority of the cells within the culture exist within spheroid or loose aggregates at the time of treatment.

For references, please see bibliography



NURTURING CELLS TO GROW

Culture media monitoring with Spedia-NMR™

The production of biologic therapeutics, such as antibodies or vaccines, requires cell cultures and fermentation bioprocesses. A full understanding of the cell media used allows scientists to first establish those factors that most impact on growth and then to determine the best way to optimise the cell upstream process. This in turn allows for the development of more effective bioprocesses, improved yields and cost efficiency, while reducing variability.

This focus on understanding and controlling the cell growth conditions as early as possible, particularly

Solid understanding of cell growth patterns and behaviours is central to the development of effective vaccines and antibodies. An innovative analysis service from Eurofins has enhanced researchers' access to the data they need to optimise research and production conditions when using cell cultures.

during the fermentation upstream development phase, led to the launch of Eurofins' Spedia-NMR™. This Nuclear Magnetic resonance-based (NMR) analysis, combined with a mass spectrometry platform method, is able to very rapidly monitor a wide selection of feed components, metabolites and toxic compounds in the culture media.

By analysing and comparing media profiles from cell cultures, Spedia-NMR™ delivers information on how cells consume and metabolise the media along the culture process. This, in turn, allows fine-tuning of the composition of the media to improve **cell viability**, reach the highest yields of biologic product and facilitate rapid process troubleshooting.

Spedia-NMR™ technology offers the biotech research and manufacturing communities significantly enhanced access to detailed quantitative data, across the broadest scope of

nutriments and metabolites. This potentially allows for a substantial reduction of the upstream process development timeline by several weeks or months, greatly improving bioreactor capacity access.

The true adaptability of Spedia-NMR™ analysis has led to its application across a range of cell systems and production platforms including bacterial fermentation, stem cells and virus production. Spedia-NMR™ is designed to perform rapid, robust and **quantitative analyses** of spent media samples in a very cost-efficient manner, using semi-automated data acquisition and processing.

the science behind

A range of NMR methods and techniques for spent cell media bioanalysis has been developed, to support the characterisation of biologics and biosimilars (including peptides, small proteins, oligo-/polysaccharides) or complex mixtures. Unlike standard methods, Spedia-NMR™ does not require systematic calibration and has a built-in matrix effect compensation. The method is not dependent on the user or machine, delivers highly reproducible data, and analyses over 50 different compounds in the media.

For references, please see bibliography



OUT OF THE BOX

PRODUCT TESTING

When nothing is available, build your own robot

The market is brimming with a vast array of consumer hygiene products, from diapers for babies to feminine sanitary items. Eurofins' sophisticated and realistic testing device revolutionised leak protection testing, providing manufacturers with reliable information about the effectiveness of products.

the science behind

The pioneering SCAN 3 simulation device can be used in the debugging phase of a product test to provide useful and reliable technical feedback, based on qualitative indicators, and so reduce R&D costs. Later in the product development process, the tool can facilitate the pre-selection of prototypes. As part of this innovation, Eurofins has developed a further enhancement to the way in which pressure is tested on products. Now completely independent of the weight of the model, pressure is still exerted from top down, but is completely variable for the first time and can be adjusted for the weight desired via a secured arm with vertical mobility.

For references, please see bibliography

In 1991, Eurofins launched SCAN 1, a first-of-its-kind testing device for diapers (nappies). This exciting dynamic "model baby" offered a valuable validation tool and the first alternative to costly and hard to implement consumer panels. Not only did it allow Eurofins to provide very real technical feedback, but also to simulate **realistic user conditions**. Since its birth, the device has evolved into a third iteration – SCAN 3.

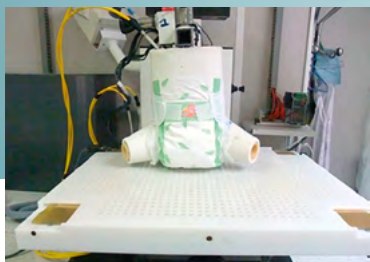
Leak protection is the best quality indicator for diapers, and this forms the core focus of the SCAN 3 device.

The parameters can be set for a wide range of realistic uses, focusing on known criteria around pressure, the quantity and number of instances of urine discharge, gender, and positions during use. Furthermore, the dynamic nature of the model's software allows testers to reproduce typical 'day-time' positions and 'night-time' positions for both sexes.

This **innovative device** is able to test not only the core absorption of the diaper but also other factors that may affect its efficiency, such as the shape of the product during wear, the degree

and manner of "gathering" during movement to a standing position, and tape resistance.

Validated since 1998, the SCAN range has seen technical advances including an anatomical ability to move the device's "legs", and the availability of devices in a range of sizes. Testers are also now able to quickly switch the device from one position to another, a feature that has further enhanced the degree of **confidence** that manufacturers, and consumers, can have in the absorbency of their products.





INVENTING THE MOST SENSITIVE HEAVY METALS DETECTION METHOD

Testing for safety in soil, water, food and pharmaceuticals

Heavy metals in foodstuffs such as fish and seafood, fruits and vegetables, and nuts and cereals are a serious threat to health. Exposure to even small amounts of cadmium, lead, mercury or arsenic can prove to be extremely toxic as they accumulate in the liver and other organs. Eurofins' pioneering tests for these substances can detect the minutest traces.

Heavy metals in the environment, in substances such as soil and water, can find their way into the food chain and pose a serious risk to consumers. The products most likely to contain these heavy metals are biological – those wholly or partly derived from biomass such as plants, trees and animals – and pharmaceuticals. Comprehensive detection of the presence of toxic materials in such products helps ensure the safety of the foodstuffs we consume and the medicines we take. By working with the latest available technology, using state-of-the-art instrumentation, and refining best practice methods, Eurofins Specialty Metals Testing, formerly known as Eurofins Frontier Global Sciences, has helped to pioneer some of the industry's **lowest detection limits** for toxic trace metals in the environment.

Operating in the cleanest possible testing environment is paramount, and the lower the detection level, the greater the importance of avoiding contamination, even at minute levels.

The US' Environmental Protection Agency (EPA) hired Eurofins as its trace metals and metals speciation reference and validation laboratory, asking the company to perform the final validation of what developed into the US EPA 1600 series methods for waters, soils/sediments and biological tissues. Today, these principles are adopted for **compliance monitoring** around the world.

Eurofins' presence and expertise in numerous other sectors meant that these methods had applications in a number of markets and industries

outside environmental testing, and the company has expanded the tests' reach to dietary supplements, the food and beverage sector, and pharmaceutical products.

Furthermore, the work of Eurofins laboratories became a catalyst for both industry and the regulatory community in **shaping policy** in several of these areas. For example, the value of differentiating different forms of arsenic in samples – such as arsenite and arsenate – helped provide the scientific means to develop policy in the food and environmental sectors due to the differing levels of toxicity of these forms of arsenic.



the science behind

The team was able to achieve lower detection limits through a combination of expert sample handling to greatly reduce laboratory-related contamination; training of customers to take samples to reduce environmental contamination; and additional proprietary measures to digest and/extract elements of concern which will allow the laboratory to use higher sampling sizes (which reduce detection limits) as well as minimise interferences. The laboratory uses a closed vessel high-heat pressurised digestion for both solids and liquids as well as microwave technology.

For references, please see bibliography

A PASSIVE REVOLUTION IN WATER SAMPLING

From spot-checking to continuous environmental monitoring

Traditional water-sampling techniques are complicated, time-consuming, and often expensive. Worst of all, they provide either only snapshots of pollution events or need expensive sampling stations that require electricity and servicing. Eurofins Sorbisense came up with a unique alternative.

Eurofins' ground-breaking Sorbisense™ **passive sampling** system is designed to ease groundwater and surface water contamination assessment, and provide better and more reliable water-quality testing data.

SorbiSystems™ comprises a small cartridge, the SorbiCell, which contains an adsorbent and a tracer salt. SorbiSystems™ is suitable for installation in groundwater, surface water and wastewater environments. Once installed, water passes through the unit at a **controlled rate**, retaining the accumulation of specific chemical groups and solutes such as pesticides, nitrate, and volatile organic compounds. These solutes are collected throughout the entire

installation period of the sampling cartridge.

As water moves through the unit, the tracer is dissolved proportionally to volume, regardless of the sampling rate. This principle allows a very accurate measurement of the sample volume and solute mass concentration. The cartridge is then easily removed and sent to a laboratory for analysis.

Data obtained through SorbiSystems™ represent an **accumulated average concentration** rather than a single snapshot, providing significantly more consistency of results and enabling an exact assessment of risks and the effect of remediation.

The technology can be used to assess groundwater, drains, drinking water, lakes, ponds and streams, and industrial wastewater. It has particular appeal for companies in sectors like aviation, oil and gas, and utilities where security issues and access to sites can be a challenge. Efficient for all types of risk screening and risk assessments, the system is estimated to save 50-80% of the previously required time in the field.

In another example of Group innovation in water sampling, Eurofins is the first laboratory network to make the **POM (polyoxymethylene) passive sampling** technology commercially available in Europe. Developed by the

Norwegian Geological Institute, POM is a passive sampling method which can be used to **monitor organic contaminants** such as PAH and PCB in water.

Thin strips of the polymer polyoxymethylene are fixed to ropes or other sampling gear and deployed from an anchor line to monitor fresh water, sea water, wastewater, groundwater and pore water, across various applications, over a four-week period or longer. POM sampling is simple to use, does not require special deployment of a field cage and avoids sample contamination or saturation.

the science behind

The Sorbisense™ sampling tool is suitable for sampling virtually all chemical solutes (contaminants). No equilibrium-based assumptions are made, and no diffusion calibrations required, just direct and transparent volume-based calculations of solute concentrations. Specially-designed mounting units allow for easy installation in a wide range of water environments.

POM is an equilibrium passive sampler where strips of polyoxymethylene are submerged in water, via an anchor line or placed in an underwater sediment bed via an aluminium frame. The POM strips monitor for organic contaminants in water over a period of four weeks or more. The state of equilibrium that the strips achieve during this time allows for the concentration in the water to be calculated, and extremely low concentrations of PAHs and PCBs can be detected. The detection limit can be as low as 1 pg/l for PCB and 1 ng/l for PAH. The testing method does not require any information on the salinity, temperature, water velocity or viscosity of the sample.

For references, please see bibliography





ALGAE, ALGAE, EVERYWHERE — TOXIC AND WHAT A STINK

First rapid test for cyanobacteria toxins in water bodies

Warm weather brings people outdoors and often to recreational waters such as beaches, lakes, ponds and rivers to enjoy those beautiful days. It can also bring about potentially toxic algae to these same bodies of water. While a number of people are familiar with "red tides," blue-green algae, more correctly known as cyanobacteria, can be just as dangerous.

Toxins produced by certain strains of cyanobacteria found in these freshwater, estuarine and marine bodies, can cause liver and kidney damage, have been linked to Alzheimer's and may be carcinogenic. Higher levels of the ratio of nitrogen to phosphorus, increased organic matter availability, higher temperatures, and decreased light intensity attenuation are among the factors that influence the formation and persistence of these cyanobacteria.

In addition, scientists have found that climate change may be a catalyst for a global increase in harmful cyanobacterial blooms. All these impacts have led to increased calls for improved testing methods for **cyanobacterial toxins**. Gold Standard Diagnostics Horsham, at the time Eurofins Abraxis, responded with the development of rapid **detection test kits** for microcystins, cylindrospermopsin, saxitoxin, and the world's first detection kits for anatoxin-a and B-Methylamino-L-alanine (BMAA).

Cyanobacteria are often confused with green algae, because both can produce dense mats that can impede activities such as swimming and fishing, and may cause odour problems and oxygen depletion which can affect fish. However, unlike cyanobacteria, green algae are not generally believed to produce toxins. Cyanobacterial blooms, on the other hand, are known to produce highly potent cyanotoxins that can damage the liver, kidneys and the reproductive system, and even cause death.

These toxins can enter the body through accidental ingestion, inhalation or dermal contact, or can end up in drinking water through the water processing system, as occurred at Lake Erie in Ohio in 2014 and which badly affected the city of Toledo. Some bacteria blooms can overwhelm the filter or capacities of water treatment plants and there is evidence that the toxin can enter the food chain via irrigation systems. Cyanobacteria do

not always produce toxins however, and the presence of toxins can only be detected through water sample analysis. The toxin microcystin was detected at Toledo by the use of kits from Gold Standard Diagnostics Horsham, part of the Eurofins network since 2017, and the test has since become EPA method 546.

Gold Standard Diagnostics Horsham is the world leader in rapid test kits for freshwater and marine toxins and continues to develop unique tests for cyanotoxins. Many of them are patent-protected and Gold Standard Diagnostics Horsham is shortly to commercialise the first ELISA – an enzyme-linked immunosorbent assay – for cyanobacterial anabaenopeptins toxins. Gold Standard Diagnostics Horsham is also developing tests to help water companies to manage cyanobacteria that produce the compounds geosmin and methyl isoborneol which have a very strong taste and, particularly, odour.



the science behind

Cyanotoxins can be produced by a wide variety of planktonic cyanobacteria. The most commonly identified cyanotoxins are microcystins, cylindrospermopsin, anatoxins, saxitoxins, and BMAA. Microcystin-LR is the most toxic form of over 160 known toxic variants, and is also the most studied by chemists, pharmacologists, biologists, and ecologists. Microcystins contain several uncommon non-proteinogenic amino acids such as dehydroalanine derivatives and the uncommon β -amino acid ADDA. Microcystins covalently bind to and inhibit protein phosphatases PP1 and PP2A and can cause non-alcoholic liver disease, pancreatitis, and liver cancer. Gold Standard Diagnostics Horsham offers two cyanobacteria toxin test formats: a quantitative ELISA which takes 1.5–3.5 hours to produce up to 40 results, and a semi-quantitative lateral flow immunoassay (dipstick) that takes around 15–30 minutes depending on the analyte.

For references, please see bibliography



The science of DNA profiling has advanced significantly over the last 20 years. The sensitivity and discriminating power of DNA tests means forensic scientists are now able to apply DNA technology in many situations that previously would have been unthinkable. However, the increased sensitivity of the technology can mean that many more 'mixed DNA profiles' are obtained. These are instances where DNA from more than one person is detected, which is common when body fluids have mixed or an object, a weapon for example, has had repeated contact from different individuals. In response to this challenge, Eurofins has developed its own proprietary software (LiRa) to assess DNA mixtures in serious crime cases.

Eurofins' proprietary DNA software warms up cold cases

SOLVING OLD CRIMES WITH NEW TECHNOLOGY



Autosomal DNA profiling was first applied in criminal investigations in the UK in the late 1980s. As DNA profiling became more sensitive, the chances of detecting DNA from multiple individuals increased and so did the incidence of mixed DNA profiles. The interpretation of some of these profiles is straightforward, while others can be very complex. Eurofins has responded by developing LiRa, its own statistical software, to help courts and investigating teams to understand the strength of evidence in cases involving complex DNA profiles. The key output from the software is a likelihood ratio (hence the name LiRa), which is calculated based on two mutually exclusive propositions defined by the user – one representing the prosecution version of events and one the defence version of events.

Depending on environmental conditions and the quality of preservation of biological material, DNA can survive for many decades. This means, provided biological material has been retained and stored correctly, crimes that took place many years before the advent of DNA

technology can be re-investigated. These crimes are referred to as 'cold cases' because they have remained unsolved despite the best efforts of investigators. Eurofins' sophisticated software has been critical to helping solve several cold cases, including a 1984 violent sexual assault in Bradford, England, for which the assailant received a conviction 33 years after the attack occurred. Forensic scientists found a tiny speck of blood on a knife recovered from the crime scene and obtained a mixed and partial DNA profile from a swab of the area. The mixed and partial nature of the result meant that computing a weight of evidence for court was complicated statistically. However, using the LiRa software, it was calculated that the DNA evidence was 490 million times more likely to be attributed to the person of interest and an unknown individual as opposed to two unknown individuals.

The software was developed, tested, and is supported on an ongoing basis by Eurofins' own in-house forensic software development team and statisticians.

the science behind

Initial DNA profiling techniques provided highly discriminating DNA profiles and compelling evidence of association, but also required several days, sometimes weeks, to produce a profile. They used radioactive probes and required a large amount of DNA (in forensic terms). Techniques have continued to evolve and become more sophisticated. Now, with the advent of PCR, DNA profiling has become super-sensitive. The old radioactive probes and Southern Blots have been replaced with fluorescent dyes and capillary electrophoresis (CE). Instead of X-ray film, DNA profiles are produced digitally and evaluated statistically, using specialist software that is costly to purchase.

For references, please see bibliography



Commercial development of genetically modified organisms (GMOs) is on the increase, with maize and soy just two examples. The emergence onto the market of the first GM animal for human consumption – a salmon – means increased calls for clarity on the sale of GM produce. Eurofins responded with the introduction of the world's first commercial test for detection of this genetically modified fish.

Fast, reliable testing for genetically modified fish

LEADING THE WORLD IN GM FISH DETECTION

GM salmon must be reared in closed fish farms without access to open water, to guarantee separation from non-GM fish. The first GM salmon, **AquAdvantage®**, found its way onto the Canadian market, and consumers' plates, in 2017, sold by producer AquaBounty Technologies. Atlantic salmon was modified with the introduction of a gene from another salmon species, the Pacific Chinook, which regulates growth hormones

allowing AquAdvantage® salmon to grow both faster and year-round instead of just during the spring and summer.

This GM salmon received approval for sale in Canada in May 2016 and, according to Canadian law, does not need to be labelled. In both 2017 and 2018, 5 tons of the GM salmon, AquAdvantage®, were sold in Canada. In 2019, the GM salmon was not only

approved for sale in the USA but also for cultivation in a salmon farm in Indiana. However, in EU member states and other countries, the marketing of AquAdvantage® salmon remained unapproved. As a result of this situation, a specific method to detect AquAdvantage® salmon was required.

Eurofins responded with the rapid introduction of a very specific and highly sensitive DNA-based test to

detect the genetic modification of AquAdvantage® salmon. The method is able to detect minute traces of the material in a sample, and to distinguish genetically modified salmon from unmodified salmon.

The method, developed by Gold Standard Diagnostics Freiburg, at the time Eurofins GeneScan Technologies, is also available as a full commercial detection kit for the modified salmon. In combination with an efficient DNA extraction method, the detection of GM material is possible even in highly processed samples. The test kit can be used flexibly with existing common DNA extraction methods and alongside one of the world's broadest and fastest-growing portfolios of existing GMO tests developed by the same company.

Gold Standard Diagnostics Freiburg in Germany is one of Eurofins' test kit competence partners and offers tests for all commercialised GMOs, including GM maize, soy, canola, rapeseed, cotton and rice.



the science behind

The Gold Standard Diagnostics Freiburg test is based on real-time polymerase chain reaction (PCR). The high specificity of PCR is achieved by the use of so-called primers and probes, short single-stranded DNA fragments, which are specific for the target DNA sequence and help amplify the target DNA sequence that is located between the two primer sequences. Whereas conventional PCR uses gel electrophoresis to make the amplified products visible, state-of-the-art real-time PCR methods use a so-called probe which is a further single-stranded DNA molecule labelled with a fluorescent reporter dye at one end and a quencher molecule at the other. The DNA polymerase, an enzyme necessary for the PCR process, degrades the probe, releasing the fluorescent reporter molecule and therefore leading to an increase in fluorescence which can be measured in real time.

For references, please see bibliography

NOT WANTED! DEAD OR ALIVE!

MOULD AND BACTERIA

Eurofins' new molecular methods enhance mould analysis

Mould and bacteria growth indoors can lead to illnesses. Work by Eurofins to improve testing methods to determine the presence of fungal and bacterial spores in indoor environments has resulted in much faster, more specific and detailed analytical tests. These tests increase the ability to detect a wider range of mould and bacteria species including those which are dead or dormant, that can develop inside buildings.

Awareness is growing of the threat to human health from mould and bacteria indoors, including **dead or dormant proliferations**. Modern buildings with higher energy efficiency may have limited air circulation and, consequently, higher humidity levels. Unfortunately, such conditions can lead to mould formation and bacterial growth. Higher numbers of these microorganisms both in air and in construction materials may affect the mucous membranes, causing respiratory ailments such as asthma, and symptoms resulting from an activation of the immune system, such as itching, eye discomfort, headaches and fatigue. In 2009, the World Health Organization (WHO) stated that mould and bacteria in indoor environments should be minimised.

Traditional testing methods, where species determination analysis is based primarily on morphology, pigmentation and growth characteristics on media, were limited to living mould and bacteria cells. In fact, dormant mould can proliferate again under the right conditions, and may also produce harmful toxins (e.g. *Streptomyces geosmin*). Traditional testing methods are also laborious and time-consuming, with samples collected either by extracting a piece of the construction material or by air sampling, before cultivating mould over seven days. Moreover, this cultivation test does not take into account any dormant/dead bacteria or mould spores.

Eurofins Environment Sweden developed a superior method to detect total mould/

bacteria biomass, so including living and dead mould. A new DNA test, developed together with Eurofins' Genomics team in Ebersberg, Germany, uses a DNA probe designed to attach to most mould found in the environment. The company also offers another similar probe for bacteria and, in addition, specific probes for some species of mould and bacteria whose odours make them undesirable. One test option produces total mould/ bacteria numbers, while another identifies bacteria and mould species by strain.

Although attribution of specific microorganisms that may occur in damaged building materials to specific illnesses remains scientifically and medically difficult, these new tests mean customers are better placed

to make informed decisions about their buildings' condition and their employees' health.

The tests can help distinguish between new ongoing moisture damage and old damage from previous mould or bacteria growths and, if total mould and/or bacteria numbers are above acceptable levels, damaged material can be removed from the building's structure and substituted with new. The tests on living and dead mould can also help evaluate different sorts of building materials for their qualities in terms of mould susceptibility.

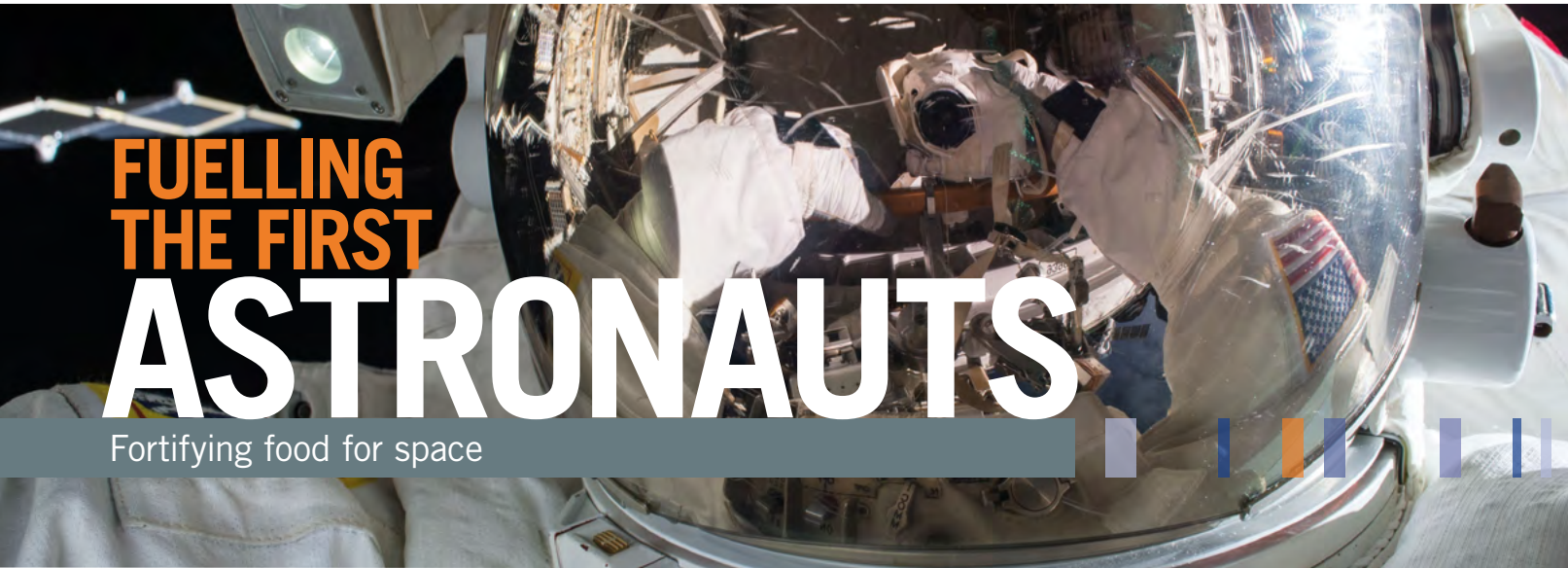
Regulatory developments in a number of countries are likely to result in the testing method becoming more widely applied.



the science behind

An ability to identify total numbers of mould and bacteria present, whether living or dead, gives scientists a much deeper understanding of the extent of bacterial presence. Applying a simple criterion that a "damaged" material contains levels of mould or bacteria more than ten-fold higher than a reference material, Eurofins researchers detected a substantial number of "damaged" samples with 10-1,000 fold higher levels than the reference sample. Bacteria were responsible for a far higher number of the "damaged" findings, and were sometimes solely responsible. The experiments also showed that living microorganism numbers were considerably lower than the total numbers, demonstrating the importance of detecting dead and dormant mould and bacteria as well as living, in order to define moisture damages causing indoor air complaints.

For references, please see bibliography

FUELLING THE FIRST ASTRONAUTS

Fortifying food for space

Have you ever wondered what astronauts eat when they are in space? With a relationship with NASA spanning over five decades, Eurofins (as Covance Food Solutions, then WARF Institute and later Hazleton) has played a crucial role in ensuring the food that fuels astronauts as they travel to outer space is nutritious, safe and palatable.

During the 1960s, The National Aeronautics and Space Administration (NASA) was heavily involved in a programme to send manned rockets into outer space, to land on the moon for the very first time. Preparation required hundreds of extremely challenging research and innovation projects, including developing the products that the astronauts would eat during the first journey to the moon.

Food scientists needed to develop fully **fortified foods**, which could remain stable for long periods of time in outer space. These foods would require lyophilisation, or freeze drying, for stability and storage, and needed to contain substantial nutrients, to sustain the astronauts, while also tasting good.

NASA partnered with Eurofins scientists, at the time WARF Institute, Hazleton Laboratories, and Covance, to determine the correct steps required to fortify these foods. The vitamins that

were to be used needed to be stabilised using an **encapsulation process**, where vitamin molecules are confined within larger molecules. Testing for these encapsulated vitamins posed a particular challenge. However, new methods were required to accurately determine how nutritious the stabilised food products were across the board, in terms of proteins, vitamins, carbohydrates, fats and minerals.

As part of the project, Eurofins scientists developed a range of **new test methods**, utilising High Performance Liquid Chromatography (HPLC) and Inductively Couple Argon Plasma Atomic Emission Spectroscopy (ICAP), and many of the methods are still in use today. Methods developed for NASA to test for vitamins and minerals in these foods have since become Official Methods of AOAC International (formerly the Association of Analytical Communities), and are used in food testing laboratories across the world.

After the extraction techniques and analyses were developed, Eurofins laboratories also put them into practice and tested all of the foods developed by NASA for their nutritional content and stability. Many of these test methods became official methods and are used today for food testing beyond the purposes of space travel.

Eurofins' close working relationship with NASA continues today as they consistently develop new, more stable, and better tasting foods for their astronauts. As NASA develops food to be used in the International Space Station, and food to be used in the first manned space mission to Mars, which must remain stable for even longer, up to a five-year period, Eurofins laboratories still perform all quality control testing on these foods used in space.

the science behind

The food products developed by NASA had to be specifically fortified with nutrients and stabilised to ensure a longer shelf life. In order to ensure that the foods still contained the correct levels of vitamins and minerals needed, Eurofins was tasked with the development of new bespoke techniques which would allow for the extraction of these vitamins and minerals to facilitate testing.

Methods used historically only had to measure natural levels of a vitamin. The NASA samples had additional vitamins added to the food, so the samples had to be treated with newly developed extraction techniques to isolate the natural vitamins and the added vitamins, a very delicate procedure.

The new analytical methods that were developed to test for vitamins in the foods created by NASA included, for example, High Performance Liquid Chromatography (HPLC) to separate, identify, and quantify each vitamin component in the foods created.

Those methods which tested for minerals made use of Inductively Couple Argon Plasma Atomic Emission Spectroscopy (ICAP-AES), a technique which is based on inductively coupled plasma to produce excited atoms and ions that emit electromagnetic radiation at wavelengths indicating a correlation to a particular element.

For references, please see bibliography

TACKLING THE BIG

Making microplastic testing commercially available

PROBLEM OF TINY PARTICLES

Microplastics are any particles of plastic fragment smaller than 5mm. Polluting our environment every day, microplastics may contain or carry hazardous chemicals that have a negative impact on nature and human health. For this reason, their ingestion through water, or foods such as fish or shellfish, is of particular concern. In 2019, the World Health Organization reviewed the state of evidence on microplastics and published a report assessing the risks to human health.

Eurofins laboratories in Norway have been at the forefront of developing testing methods to identify and quantify microplastics in the environment using **pyrolysis-GC/MS**. By their very nature, microplastic particles are just that – micro. Microplastic particles can be filtered

from clean liquid samples; and for more complicated matrices, a sample treatment step must be performed prior to analysis. The team has developed methods for the analysis of microplastics in various liquid matrices, wastewater, sediments and soil, fish and dry matter for polymer screening. Rubber components, mostly found in tyre wear particles, can also be analysed using **pyrolysis-GC/MS** which can provide important information about the pollution in the environment from traffic.

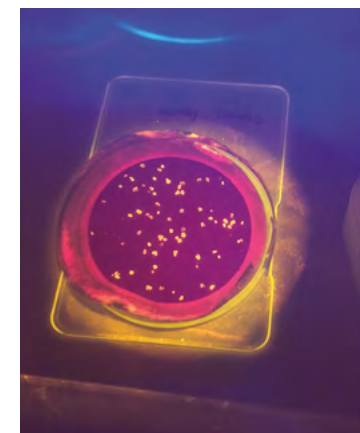
Several methods can be used to identify individual microplastic polymers – the most common being dye imaging and NOAA methodology under **FTIR spectroscopy** and microscopy technologies. Eurofins SFA Laboratories, based in Wisconsin, was the first

laboratory to make these analytical methods commercially available, offering testing across several matrices including marine and drinking water sources and finished bottled water. Additionally, both Eurofins Analytical Services Hungary Kft. and Eurofins Environment Testing Australia offer state-of-the-art spectroscopic analysis for microplastics in a diverse array of products such as infant formula, eye drops, soy sauce, rice, sea salt, shellfish and anything where plastic has been used in the processing and/or packaging.

Eurofins Textile Testing Spain has been one of the first testing laboratories to develop a quantitative detection method to monitor **microfibre-shedding during the washing process** of synthetic textiles such as polyester, acrylic, elastane, and nylon. These test

If not correctly recycled, plastic poses a major threat to our environment. Degrading into tiny particles, microplastics can remain in the environment for over hundreds, if not thousands, of years, infiltrating aquatic and marine ecosystems and contaminating the water we drink. Recognising the major global environmental problem that microplastics present, Eurofins is at the forefront of providing testing solutions to identify the prevalence of microplastics in our environment.

methods help customers to better understand the environmental impact of their products, and by identifying which fabrics and fibres are most prone to microfibre release, enables them to proactively reduce microplastic pollution through informed raw material selection and use. More recently, Eurofins laboratories from several countries have been involved in international research to measure airborne microplastic particles. Samples were taken in the same way (for the same duration, using the same method, etc.) at several locations around the world for testing by experts in the various laboratories. Across the network, Eurofins companies continue to undertake research to better understand the threat that microplastics pose to our environment and their prevalence in our ecosystems.



the science behind

FTIR spectroscopy and microscopy analyses use infrared light to scan test samples and observe chemical properties. In the dye imaging method, microplastic particles in water absorb dye and glow under ultraviolet light, highlighting their presence. The NOAA Method identifies larger microplastic particles in water which is considered to be dirtier. The water undergoes chemical treatment, density separation, mass determination and possible spectroscopic investigation.

Pyrolysis-GC-MS testing utilises heat to decompose samples to produce smaller molecules which are then separated by gas chromatography and detected using mass spectrometry. Pyrolysis-GC-MS can be applied to insoluble and complex materials including plastics, and may also be combined with FTIR spectroscopy to provide number of particles, characterisation of polymers and weight.

For references, please see bibliography



SAFEGUARDING THE HEALTH OF THE WORLD'S MOST

Setting the new standard in infant formula testing

VULNERABLE CONSUMERS – INFANTS

In 2008, China was rocked by a national food safety scandal when it emerged that a chemical component called melamine, which is harmful to humans and is used in plastic and fertiliser production, had made its way into infant formula and milk. Of an estimated 300,000 victims affected by the compromised milk and infant formula, six babies died and an estimated 54,000 babies were hospitalised. The tragedy highlighted the importance of using the most stringent methods to test infant formula. Having screened tens of thousands of samples for melamine levels during the Chinese scandal, Eurofins companies have also played a crucial role in a global initiative to overhaul, redevelop and modernise infant formula testing.

Infant formula is often the sole source of nutrition for infants and must contain 100% of their required nutrients. By its very nature, it is one of the most highly regulated products in the world, and it is essential that the most stringent validated testing methods are used for product analysis. The 2008 Chinese milk scandal was described by the World Health Organization as one of the largest food safety events it had ever had to deal with, and noted that the crisis of confidence among Chinese consumers would be hard to overcome. At that time, the standard test methods for infant formula were those developed by AOAC International (formerly the Association of Analytical Communities) in the 1980s, which had become outdated. The scandal was the catalyst for an initiative to modernise and improve all infant formula testing methods.

As members of AOAC, leading infant formula producers such as Abbott, Danone, Johnson & Johnson and Nestlé came together in 2010 to form



the Stakeholder Panel on Infant Formula and Adult Nutritionals, **SPIFAN**, to develop a new uniform suite of stringent infant formula testing methods. Scientists from Eurofins Food Chemistry Testing Madison and Botanical Testing Brea CA, at that time Covance Food Solutions, were chosen to lead SPIFAN, as Chair of the stakeholder panel, and were chosen to validate all new testing methods. Eurofins scientists also held positions as Working Group and Expert Panel Review Chairs, as well as method developers and reviewers.

SPIFAN identified and validated a modern suite of best practice analytical test methods, the current AOAC Official methods for nutrition in infant formula and adult nutritionals. As a result, 65 new AOAC Official Methods have been validated by SPIFAN, which are regarded as the world and industry standard in infant formula testing.

New methods developed to test for fat soluble vitamins, A, D and E for example, are considered to be among the most accurate and precise methods ever developed.

Eurofins scientists participated in the collaborative study trials for over 15 of these new SPIFAN methods. The new approved SPIFAN methods for the analysis of Vitamin D, Iodine, Choline, and Carnitine were all developed and validated by these scientists, and are considered the international 'gold standard' in testing of this kind.

The redeveloped tests which are based on different methods than those used in the 1980s, are nowadays applied by every infant formula producer across the globe. They are recognised by international standards groups such as the International Standards Organization (ISO), the International Dairy Federation (IDF), and Codex Alimentarius. SPIFAN, and the test methods developed by Eurofins, changed the landscape of infant formula testing, and showcased Eurofins' ability to create world-class testing standards.

Eurofins Expertise Centre for Complex Carbohydrates co-developed, in close cooperation with Nestlé Research,

a novel method to determine all inulin-type fructans found in infant formula and adult nutritional products. In addition, the Group established a new standard method that allows for testing for 2'-fucosyllactose (2'-FL), the most abundant of the human milk oligosaccharides (HMOs). Several studies have shown the importance of HMO for a baby's development.

Most recently, Eurofins Food Integrity and Innovation (EFII), USA, developed an optimised United States Food and Drug Administration (FDA) **method for the analysis of 16 target Perfluoroalkyl Substances (PFAS)**, validated for infant formula and related ingredients. Eurofins LabCo in Rotterdam has also launched a **fatty acid composition analysis** that is approved by AOAC's SPIFAN. In this method, sample preparation is integrated and therefore variation is reduced. This method helps infant formula and dairy producers to optimally dose ingredients to deliver a higher quality product.

the science behind

The Vitamin D, Iodine, Choline, and Carnitine methods developed by Eurofins made use of some of the most advanced analytical instruments at the time.

The assays for Vitamin D, Choline and Carnitine involved the use of LC MS-MS (Liquid Chromatography Tandem Mass Spectrometry) in vitamin analyses for the first time. This is an analytical chemistry technique that combines the physical separation capabilities of liquid chromatography with the mass analysis capabilities of mass spectrometry, far more accurate, precise and specific than the HPLC, colorimetric and ELISA methods used pre-SPIFAN.

The Iodine assay redeveloped by Eurofins utilised ICP-MS, where equivalent tests previously used ion specific electrode testing. ICP-MS, or inductively coupled plasma mass spectrometry, is capable of detecting metals and several non-metals at very low concentrations. This is achieved by ionising a sample with inductively coupled plasma and then using a mass spectrometer to separate and quantify ions.

The fructan analysis method has been designed to determine fructan content without the need for ingredient-specific correction factors, thus avoiding the need for pre-analysis or prior knowledge of the fructan type in the product. This method combines enzymatic and SPE steps to remove interfering components.

The new sugar analysis method was designed to overcome issues in relation to interference of other saccharides in infant formula like lactulose, HMOs or fructans. Finally, the new method to test for 2'-fucosyllactose (2'-FL) allows for the first time the detection of this important HMO.

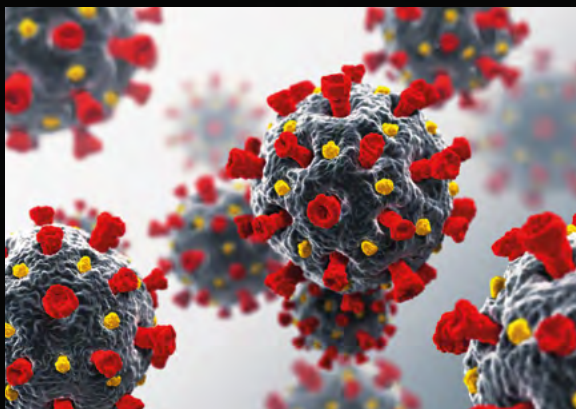
For references, please see bibliography



20 MILLION TESTS A MONTH TO HELP PROTECT THE WORLD AGAINST

Innovation and action during a global pandemic

COVID-19



In 2020, Eurofins' laboratories and colleagues pulled together to contribute to the ever-changing global efforts to safeguard global health and societies during the COVID-19 pandemic, the effects of which reached every corner of the world. Widespread testing was a cornerstone of the strategies developed by governments and national health authorities across the globe to protect their societies from the virus. The World Health Organization declared COVID-19 to be a pandemic in March 2020, and by May 2020, Eurofins companies had already created a large range of products and services to facilitate over 20 million patient tests per month globally. Eurofins' support included optimised SARS-CoV-2 (the virus that causes COVID-19) specific probes and primers, used by a number of independent kit producers.

Eurofins teams worked round the clock to establish and **ramp up COVID-19 clinical testing capacity** to identify

In 2020, the world was rocked by COVID-19, a potentially life-threatening and highly infectious respiratory disease, caused by a novel coronavirus. COVID-19 was labelled a global pandemic by the World Health Organization in March 2020. In the immediate outbreak, countries across the world grappled with the devastating effects of the virus on their citizens, public health systems and economies. Eurofins acted quickly to develop and ramp up testing capacity to support governments to safeguard public health and innovated to bring urgently required tests to the market to support the fight against COVID-19.

those carrying the virus, using the real time reverse transcriptase polymerase chain reaction (RT-PCR) method, and returning results within 12-24 hours. This meant expanding Eurofins laboratories' testing capacities, but also developing new products that would allow operators to perform the actual testing (nucleic acid extraction kits, RT PCR kits, etc.).

As the pandemic progressed, **antibody testing** played an increasingly helpful role, by identifying people who may have been exposed to the virus, possibly without even noticing, and allowing health authorities to evaluate population exposure and immune response. This provided insights to inform policy decisions such as vaccination strategies for example, by indicating the most vulnerable groups, and, at a time when vaccines were not very widely available, determining which groups should receive the vaccine first. Eurofins developed antibody testing services and kits so

that those who may have been sick, but could not be tested at the time could determine if they already had antibodies as a result of exposure to COVID-19. Eurofins launched particularly fast rapid point-of-care finger-prick testing devices which identified antibodies in just 10 minutes, with a sensitivity of 94.5% from 19 days following the onset of symptoms.

Eurofins also developed **T-cell assays**, which facilitate the detection of the T-cells specific to the virus in a patient. This is important because variations in T-cell populations have been reported to provide insights about the outcome of disease progression, and these help doctors to plan for ICU bed and ventilator capacity, especially in times where there may have been shortages.

During the initial outbreak of the virus, governments across the world chose to "lock down" economies and societies. As economic and social activity returned, it was obvious that ongoing surveillance

would be necessary to quickly identify any resurgence in COVID-19 cases and monitor the spread of the disease across all types of environments and workplaces. Eurofins developed **monitoring methods and kits** such as environmental surface testing, wastewater testing, air testing and worn mask testing methods that could detect the presence of SARS-CoV-2 at very early stages, and, as a result, minimise spread.

Eurofins research showed that workplaces with SARS-CoV-2 contaminated surfaces were 10 times more likely to have COVID-19 positive employees than locations with no or very few contaminated surfaces. Environmental **surface testing** methods allow for early warning of potential virus outbreaks by detecting the presence of pre- or asymptomatic virus spreaders at a site or in an environment without the need for invasive and costly human clinical testing. Asymptomatic people are persons infected with the virus,

but without symptoms of disease, and pre-symptomatic are those who are harbouring the disease but have not yet shown symptoms. Both may unknowingly transmit viruses to others.

Eurofins was the first commercial laboratory in Europe to offer COVID-19 wastewater testing. **Testing wastewater** can provide an early indication of the presence of viruses in a specific community or at a work site. Faeces from people infected with the SARSCoV-2 virus can contain genetic material (RNA) from the virus, and that viral RNA can be detected in sewage and wastewater. Studies have shown wastewater testing capable of detecting a community COVID-19 prevalence rate as low as a 0.02%- 0.1% (i.e. between 2 virus shedders per 10,000 persons and 1 virus shedder per 1,000 persons). In Denmark, for example, Eurofins' wastewater testing method detected SARS-CoV-2 in wastewater two days before the first official COVID-19 case was confirmed in the country. This





the science behind

Real time reverse transcriptase polymerase chain reaction (RT-PCR), the method used in COVID-19 detection tests, is a laboratory technique which combines reverse transcription of RNA into DNA and subsequent amplification of the molecules. Thereby minute traces of virus material can be detected. RNA is the molecule the SARS-CoV-2 virus uses as genetic information. New variant screening worked by detecting single nucleotide polymorphism (SNP), a variation at a single position in the viral RNA sequence, to identify the variant responsible for the infection.

The unique fragment length analysis (FLA) SARS-CoV-2 detection kit utilises Sanger sequencing, an alternative to RT-PCR. The method detects the reverse transcribed, amplified viral RNA on Sanger capillary sequencers and can help to ramp-up testing capacities.

The serological test, a blood test that looks for antibodies, detects infection after the immune system has produced antibodies (IgA, IgG or IgM) against the virus, which remain in the bloodstream for some time after the patient has recovered.

For references, please see bibliography

technology was also successfully used to detect asymptomatic spreaders on college campuses. Wastewater testing in sewage plants is commonplace in many countries around the world, and so this method could be easily rolled out and provide crucial information to communities. Like environmental surface testing, wastewater testing allows for the identification of asymptomatic spreaders, and supports organisations to monitor the safety of their environments. Eurofins also made its wastewater testing kits available for purchase to other laboratories as well as state and public health authorities.

Environmental surface, air and worn mask testing as well as wastewater monitoring allowed for human clinical testing to be focused where virus presence was likelier, allowing for the allocation of capacity-constrained human COVID-19 testing where it was most needed.

Besides carrying out testing in their own laboratories, several Eurofins companies received approval for their laboratory-developed tests to be used for the detection of SARS-CoV-2 in patients by public and private laboratories. Eurofins also developed dedicated reagents to support the entire testing process: from nucleic acid extraction kits, to PCR kits for SARS-CoV-2 detection and for specific variant identification, to kits for SARS-CoV-2 sequencing, antibody testing and T-cell monitoring, including equipment and point of care solutions. Eurofins sites created capacity to provide primers and probes to be incorporated in millions of SARS-CoV-2 **testing kits** worldwide. Besides developing kits for Eurofins and other laboratories to carry out PCR testing, Eurofins also developed a unique SARS-CoV-2 detection kit utilising sequencing equipment common in molecular testing and Genomics laboratories. This method, which is as sensitive as PCR testing, resulted in much needed additional testing capacity.

During the pandemic, Eurofins companies also **facilitated the sequencing** of SARS-CoV-2 to identify new virus strains or Variants of Concern as they emerged. The company opened a high-throughput sequencing laboratory to analyse the full genome of SARS-CoV-2 and support the European Center for Disease Control. Between February 2021 and February 2022 more than 250,000 viral genomes, extracted from samples collected in 24 countries, were fully sequenced and analysed thanks to this collaboration, contributing to the identification and tracking of emerging virus strains.

Early detection of particularly virulent variants with increased transmissibility was key to delaying their introduction and spread within the population. With this objective in mind, Eurofins companies developed more than 15 CE marked clinical kits and more than eight water surveillance and surface tests for variant screening, all of which were

brought to the market in record time. On the day that Omicron was officially designated a Variant of Concern by WHO, Eurofins companies launched a screening solution to identify the variant.

Eurofins laboratories also carried out emergency testing of medical devices, such as respirators, and **Personal Protective Equipment (PPE)** during the pandemic.

Eurofins' unmatched global network of BioPharma Product Testing laboratories ramped up capacity to support some of the largest global pharmaceutical and biotechnology companies developing products to fight COVID-19 and launched products and services aimed at supporting the research necessary to **develop novel vaccines and therapeutics**.

PROTECTING CROPS AND HUMAN HEALTH AT THE SAME TIME

Developing the first test kit to detect glyphosate

Human exposure to herbicides through food consumption can pose potential health risks. According to a 2016 study, glyphosate is the most common agricultural herbicide in the world, used to kill weeds that compete with crops. Gold Standard Diagnostics Horsham, at the time Eurofins Abraxis, developed and brought the first rapid test solutions, from semi-quantitative field screening to quantitative and automated laboratory testing, to detect glyphosate, to market.

Food consumption is an important pathway of human exposure to pesticides, herbicides and chemical contaminants such as glyphosate. Herbicides play an important role in improving crop yields and feeding a growing population, but the responsible use of these items is in the best interests of humankind and the environment. Glyphosate and other chemical compounds can enter the food supply through direct application to food crops, indirect application by drift from aerial spraying of adjacent fields, through soil or through cross-contamination of shared equipment during food processing.

Glyphosate is absorbed through foliage and transported to growing parts of a plant. As such, it is only effective on actively growing plants and not as a pre-emergence herbicide. A great number of crops, such as soybean and corn for example, have been genetically engineered to withstand glyphosate, and so farmers use glyphosate as a post-emergence herbicide against weeds.

While glyphosate has been approved by regulatory bodies worldwide, concerns about its potential effect on humans and the environment persist. As a result, **consumer demand** for

"glyphosate free" food products has been growing for several years. Leading food manufacturers demanded that suppliers decrease the concentration of glyphosate in the grains they produce, and consumer pressure has driven manufacturers to carry out glyphosate testing on their food products.

Gold Standard Diagnostics Horsham, at the time Eurofins Abraxis, developed the first commercially available **rapid test kit**. The kit detects low levels of glyphosate in crops safeguarding consumers and supporting manufacturers to identify and producers to showcase "low glyphosate" crops.

Gold Standard Diagnostics Horsham's Abraxis® Glyphosate ELISA (enzyme-linked immunosorbent assay) and lateral flow (dipstick) test kits allow for the sensitive (low ppb, parts per billion) and rapid analysis of glyphosate in grains, coffee, and over 30 other matrices. Lateral flow tests can be carried out rapidly in field, while ELISA test kits are subject to traditional laboratory testing.



the science behind

Specific antibodies are required for ELISA to be successful. As glyphosate is a very small molecule, it is very difficult to develop an antibody against it. Eurofins managed to do so, and thus developed both of its immunoassay-based glyphosate detection tests as a result. The ELISA test utilises microtiter plates for a fully quantitative analysis using a set of standards, while the lateral flow test strip provides a qualitative/semi-quantitative result around specific test levels.

ELISA is a plate-based assay technique designed to detect and quantify soluble substances such as peptides, proteins, antibodies, and hormones.

The lateral flow test strip is a diagnostic device which is used to confirm the presence or absence of a contaminant or other target analyte in water supplies, foodstuffs, or animal feeds. The test is semi-quantitative at target levels (e.g. 40 ppb; 10 ppb).

For references, please see bibliography



WHAT DO PIZZA BOXES, RAINCOATS AND FRYING PANS

Detecting PFAS in everyday products and our environment

HAVE IN COMMON?



PFAS comprise a group of thousands of synthetic chemicals that have been in commercial production since the 1940s. They can be found in everyday products like non-stick cookware, cosmetics, fire retardants, stains, water repellents, some furniture, waterproof clothes, pizza boxes, food packaging, carpets and textiles, firefighting foam, rubbers and plastics, electronics, and even some dental floss. The primary sources of PFAS in the environment are manufacturing facilities and PFAS-containing products that have gone into landfills over the decades, as well as firefighting foams. PFAS are largely unsusceptible to degradation, remain in the environment almost indefinitely, and are often termed "forever chemicals". Their water solubility allows them to penetrate soil to enter ecosystems.

Some PFAS accumulate over time to increased concentrations in humans, animals, and the environment. They are dispersed through air and water, to such an extent that they have even been found in the environment of the Arctic and its polar bears, and open ocean

waters. PFAS have also been found in fish, shellfish, vegetables, and other produce grown in contaminated soil or water. The Centers for Disease Control and Prevention (CDC) found PFOA, a specific PFAS, in the blood of 98% of Americans, as well as in breast milk and umbilical cord blood.

The health effects of the various kinds of PFAS are under constant research, however evidence has linked exposure to some of the chemicals to developmental issues, cancer, liver damage, immune system disruption, resistance to vaccines, thyroid diseases, impaired fertility and high cholesterol. Certain PFAS have been classified as "possibly carcinogenic" to humans by the US Environmental Protection Agency (EPA) and the International Agency for Cancer Research (IARC).

With public concern regarding the health effects of PFAS outpacing the regulatory framework to address these complex chemicals, the scientific community stepped in and worked to develop an approach to assess the

PFAS, per and polyfluoroalkyl substances, are a class of synthetic chemicals that you may not have heard of, but have certainly been in contact with. These chemicals have unique and useful characteristics and can be used to make products heat and stain resistant, non-stick, and water repellent. They are present in a myriad of consumer products, from the pans we use to cook our food, to our raincoats, and even some dental floss. Unfortunately, they are very stable compounds that accumulate and persist in our environment and have been linked to a number of health risks. Eurofins companies have been at the forefront of developing methods to test for these compounds.



prevalence of PFAS in the environment. Eurofins Environment USA became **the first commercial laboratory to test environmental samples for PFAS** in the USA. In 2016, Eurofins Environment Testing Sweden AB became the first accredited commercial laboratory to be able to detect PFAS in water at LOQs (limit of quantification) of 0.3 parts per trillion (ppt). One ppt is equivalent to a single grain of sand in an Olympic sized swimming pool. These levels are even lower than those required by the European Union Water Framework Directive (WFD). PFAS testing is a field where change and learnings are developing rapidly, and being at the frontier of research is extremely important.

Eurofins Environment Testing Sweden was also the first major commercial laboratory to introduce an analysis for a new class of PFAS in water for so-called "ultrashort PFAS". One such "ultrashort PFAS", for instance, is Trifluoromethanesulfonic acid (TFMS), which is used in Lithium-ion batteries and has already been detected in drinking water in Germany.

Of the estimated thousands of PFAS chemicals, laboratories are only able to analyse for about 100 known PFAS compounds, as many remain largely unidentified or lack analytical reference standards. Eurofins' teams of expert scientists and its investment in state-of-the-art equipment has made identifying some of these unknown PFAS in the environment and their sources possible. In 2016, Eurofins introduced the **Total Oxidizable Precursor Assay** technique, which allows for the measurement of unknown PFAS by identifying precursors of PFAS that have the potential to transform, when oxidation occurs in the environment, into the PFAS compounds that can be analysed. Eurofins Environment USA has been called upon to support the validation of several US EPA methods to measure PFAS in potable and non-potable water as well as solids.

In addition to **water** testing, Eurofins companies have also expanded their PFAS testing across other matrices, including **food** and **feed**. In 2023, e.g., the EU set limit values for PFAS in

food like fish, meat, and eggs, and the importance for testing food for these chemicals is growing. Also in 2023, Eurofins Consumer Product Testing and Eurofins Sustainability Services' global network of laboratories launched Total Fluorine (TF) and Total Inorganic Fluorine (TOF) PFAS screening services for textiles and other consumer products' matrices to augment their targeted PFAS testing. Eurofins Medical Device Services USA is pioneering PFAS testing in the medical device industry as one of the first laboratories to offer GMP PFAS testing and PFAS Risk Assessments in 2025.

Additionally, through a Eurofins empowerDX and Eurofins Environment Testing collaboration, the companies launched the first direct-to-consumer PFAS Exposure™ Test to identify the chemicals in blood samples. As a result, Eurofins Belgium was tasked in 2022 to carry out the largest European biomonitoring project for PFAS in blood, commissioned and funded by the Agency for Care and Health of the Flemish Government. The blood of over

75,000 inhabitants within a 5 km zone around the 3M factory in Antwerp will be analysed for the presence of PFAS.

Today, a growing concern regarding PFAS is how they are spread through air pathways. PFAS are emitted into the air from manufacturing facilities or facilities that treat waste of products that comprise PFAS. Eurofins companies have pioneered sampling and analytical techniques to collect and analyse PFAS from these sources, including soil vapour and stack emissions, and monitor ambient **outdoor and indoor air** for these chemicals to collect the information we need to keep the air around us safer. Such specific analyses require unique and highly specialised equipment, often only found in research laboratories and academia. Eurofins companies maintain more of this specialised equipment than any commercial laboratory in the USA and are using it to develop techniques with the aim of closing some of the data gaps that currently exist in relation to PFAS.



the science behind

The Total Oxidizable Precursor Assay technique is a chemical oxidation of PFAS precursors yielding measurable perfluoroalkyl acids. The aim is to assess the "unknown pool" of compounds that, with time, may form other, more stable PFAS in the environment.

Eurofins Environment Testing laboratories developed LC-MS/MS analysis (Liquid Chromatography with tandem mass spectrometry) for several classes of PFAS and have expanded both the list of PFAS and matrices analysed. These matrices include groundwater, soil and sediment, tissues, serum and foodstuffs, but have recently expanded to textiles and other consumer products and even the air one breathes. LC-MS/MS is a powerful analytical technique that combines the separating power of liquid chromatography with the highly sensitive and selective mass analysis capability of triple quadrupole mass spectrometry. With these tools, Eurofins achieves industry leading detection limits: as low as parts-per-trillion (ppt) detection limits for aqueous matrices and less than a part-per-billion (ppb) for solids and tissues.

For references, please see bibliography



PROTECTING CANCER PATIENTS

Identifying high-risk patients following stem cell transplant

For many cancer patients, the transplant of stem cells from a donor is potentially a life-saving treatment. However, there is a risk that donor cells can attack the patient receiving them in a disease called acute graft versus host disease (aGVHD), a potentially life-threatening condition following cell transplantation. Innovative tests, offered by Eurofins companies can predict the risk of rejection so that the best patient care can be planned for.

Stem cell transplants, including peripheral blood, bone marrow, and cord blood transplants, can be used to treat cancer, most often those affecting blood or the immune system, like leukaemia, lymphoma, or multiple myeloma. Bone marrow and stem cell transplant treatments use very high doses of chemotherapy, sometimes coupled with radiotherapy, to try to eliminate cancer cells. While the high dose treatment can kill cancer cells, it also can remove the stem cells in bone marrow that produce blood cells. Soon

after treatment, stem cells from a donor are given to a patient to replace those that were destroyed. The donor stem cells make their way into bone marrow and start to produce healthy blood cells again. A bone marrow or stem cell transplant from another person, either from a matched relative or an unrelated donor, is called allogeneic hematopoietic stem cell transplantation (HSCT).

While potentially life-saving, allogeneic HSCT can also lead to the

development of **acute graft versus host disease** (aGVHD). aGVHD is caused by immune dysregulation that is initiated when particular types of allogeneic white blood donor cells (the graft) react against the cells of the patient (the host) receiving them. The donated cells recognise the host as foreign and attack, damaging host tissue, and releasing inflammation fostering substances into the body. aGVHD typically occurs in the first three months post-transplant, has an incidence rate of between 19 – 66% and can be life threatening in severe cases.

Work over the past several years has investigated the use of **biomarkers** for non-invasive and predictive assessment of aGVHD risk. Biomarkers can determine whether aGVHD is likely to develop so that clinicians can accurately identify, at an early stage, patients which are at high-risk for severe aGVHD and improve patient outcomes following allogeneic transplants. In August 2018, Viracor Eurofins became the first laboratory

to make a predictive aGVHD assay commercially available.

The assay predicts the risk of aGVHD development in those patients who are beginning to display symptoms, those who have previously received aGVHD treatment and even those who are not yet presenting any symptoms indicating the disease. The scope of the test, made available by Eurofins companies supplies healthcare providers with actionable data to definitively **identify at-risk patients** and guide decisions to improve treatment outcomes. High-risk patients can be preemptively treated for aGVHD, while low-risk patients may potentially avoid unnecessary treatment. The aGVHD predictive assay provides a faster, more cost effective and less invasive testing alternative to biopsy, and provides vital information to allow for preemptive intervention, minimising the risk of relapse and infectious complications.



the science behind

The aGVHD predictive assay was developed by Drs. James Ferrara and John Levine (both professors of Pediatrics, Medicine, Haematology and Medical Oncology at The Tisch Cancer Institute at Icahn School of Medicine at Mount Sinai), and validated in conjunction with 17 hematopoietic stem cell transplantation (HSCT) centres to help predict the risk of non-relapse mortality (NRM) and aGVHD in HSCT patients.

The assay is based on algorithms utilising serum levels of the so-called ST2 and REG3 α biomarkers. These algorithms have clinically validated cutoffs that provide actionable insights within 24 hours. High-risk patients can be preemptively treated for aGVHD by adjusting immunosuppressive drug dosages prior to the onset of clinical disease, while low-risk patients may potentially avoid unnecessary treatment.

For references, please see bibliography

SEEKING OUT

Developing the first test to identify Radon

A SILENT KILLER

Exposure to Radon, an odourless, colourless gas, is a health hazard, and is responsible for more deaths in Sweden each year than road traffic accidents and 14% of all lung cancer diagnoses in the country. Eurofins (as MRM Konsult) commercialised the first method to test for Radon in air, which is still used all over the world today, to protect societies from this dangerous gas.



Radon is a naturally occurring radioactive gas that is produced when uranium in rocks and soil decays. It can only be identified using specialised equipment and test methods. In the open-air, the gas is diluted to harmless levels. But in enclosed spaces it can build up to dangerous amounts and become a health hazard. Radon from the ground can enter buildings and houses through cracks in floors or gaps around piping or cables. Radon poisoning does not cause obvious symptoms often associated with radioactive substances, but rather long-term exposure can lead to the development of lung cancer.

Based in Luleå, Sweden, Eurofins, as MRM Konsult, was the first laboratory in the world to make testing for Radon in air commercially available. A method to **detect Radon in indoor air**, known as the "track etch" method was first developed in the 1990s. Well regarded for its reliability, the method, which has now become the **ISO Standard 11665-4**, is a global reference and used by laboratories across the world.

The test is carried out by placing a detector in a building for approximately two months. As air diffuses through the detector, any amount of Radon in the air is measured. The detector contains a special plastic film which is damaged when Radon is present in the air. The lesions on the film are then analysed to assess the levels of Radon present.

Other than in indoor air, Eurofins is also testing for Radon directly in soil, using a method based on measuring radioactivity with active charcoal.

In 2013, the European Union introduced regulation requiring workplaces to monitor for the presence and analyse the level of Radon in the indoor environments where their employees work. Eurofins Radon Testing Sweden conducts approximately 15% of all Radon measurement tests carried out across the world.



the science behind

The method to analyse Radon in an indoor environment is based on passive sampling (using a collecting medium to accumulate chemical pollutants in the environment over a longer period of time) and uses integrated measurement equipment and deferred analysis to determine the average active concentration of Radon in a space. Eurofins' passive Radon detector has a broad measuring range. The detector can be used for long-term radon measurements or short-term screenings. It contains a plastic film and a lid made of electronically conductive plastic. When Radon enters the device, it decays into its "daughter products" also known as "decay products", thereby releasing alpha radiation and hence energy, and this energy causes damage to the plastic film. The lesions on the film are then assessed by the amount of damage per cm²; and their shape, depth and size.

For references, please see bibliography



ENSURING LIQUID GOLD IS THE

Innovative methods for fraud detection in honey

REAL DEAL



As consumer demand for natural sweetening products like honey continues to increase, so too do instances of fraud or fraud attempts by dishonest suppliers. In Europe, for instance, the European-Commission-led initiative "From the Hives" confirmed that a significant percentage of imported honey products do not comply with EU purity benchmarks. To protect consumers and legitimate domestic producers, Eurofins companies have developed a suite of state-of-the-art authentication methods to distinguish syrup-based alternatives from genuine honey.



Honey is widely regarded as one of the most **adulterated** foods in the world and testing to determine its authenticity is therefore extremely important. Authorities across the globe are facing rising fraud attempts, this can be seen in New Zealand through the first "fake Manuka honey" prosecution trial in 2019, or the action of the European Union to launch its "From the Hives" initiative in 2021, to assess the prevalence of honey adulterated with sugars on the EU market.

The composition and characteristics of honeys are subject to broad variation due to the numerous honey varieties and origins worldwide. This makes the reliable and decisive detection of possible fraudulent sugar syrup addition particularly difficult. In addition, fraudsters continue to push limits, and refined syrups, which often perfectly mimic honey composition, are also added by producers. These factors mean that particularly sensitive and reliable complementary testing methods are required to protect brands and customers. Eurofins has established

the most advanced and reliable testing method for honey authenticity.

In 2018, Eurofins Food Integrity Control Services GmbH was the first laboratory worldwide to offer authenticity analysis of honey by liquid chromatography coupled with high resolution mass spectrometry (LC-HRMS) as an ISO 17025 accredited method. The method enables the detection of known as well as unknown honey adulterants (targeted and untargeted analysis) as part of just one analytical test and has a significantly higher sensitivity for foreign sugar adulterants compared to previously established honey authenticity testing methods. The method combines a number of single detection methods that test for specific additives or adulterations into one multi-method which can **simultaneously detect different types of sugar syrups** used for adulteration, making testing more efficient and cost-effective. The Eurofins laboratory was the first laboratory worldwide to have developed a new LC-HRMS method for verification of the country of origin of honey. This method builds on the

Group's established reputation as a leader in authenticity testing for honey.

In 2014, the Eurofins Laboratory in Nantes had already pioneered a new analytical approach to test honey integrity. Developed as part of a collaborative research project, this holistic method, which uses high resolution Nuclear Magnetic Resonance (NMR), provides a wide range of information that is both targeted (quantification of defined substances) and non-targeted (identifying deviations from reference spectra). It can simultaneously detect the **addition of sugars from any source**, or other irregularities in the honey, such as excessive heat treatment or fermentation, and also confirm the botanical and geographical origin, independently from conventional pollen microscope analysis. An example of such advanced origin control is the control of Manuka honey authenticity by NMR. Eurofins' optimal offer for honey is a combination of these advanced methods with essential conventional ones such as pollen microscopic analysis.



the science behind

The LC-HRMS testing method is an analytical chemistry technique that combines the physical separation capabilities of liquid chromatography with the capabilities of high resolution mass spectrometry to analyse and quantify compounds, determine elemental compositions, and identify unknowns. Given the extreme sensitivity of the LC-HRMS testing method, it can largely narrow down possible chemical formulas of the unknown compounds. LC-HRMS can be used as a multimethod with hundreds of identified marker substances for the simultaneous detection of additions of different types of sugar syrups to honey. Combined with ¹H NMR-Profiling, ¹³C stable isotope analysis and pollen analysis, it is the most advanced and reliable testing method for honey authenticity.

NMR profiling is a technique used to identify the characteristics of food – its authenticity, origin and stability – by determining specific individual profiles for food product types, like fingerprints. Eurofins carries out NMR profiling based on its worldwide database of honey (fingerprints), which comprises over 30,000 reference samples, from over 130 different botanical families. These reference samples have been collected from local producers in more than 65 countries across the world.

For references, please see bibliography



DETECTING

Optimising outcomes for kidney transplant patients

A SILENT THREAT – TRANSPLANT REJECTION



Kidney transplantation is universally recognised as the best treatment option for patients with end-stage renal disease. Although clinical outcomes of kidney transplantation have improved steadily for decades, transplant rejection remains one of the key challenges to long-term patient survival. Eurofins' TruGraf® test is the first test capable of reliably ruling out "silent" subclinical acute rejection in patients with stable renal function, where a patient's immune system can reject a kidney transplant without the patient showing any other clinical symptoms.



According to the Global Observatory on Donation and Transplantation, approximately 90,000 kidney transplants are being performed across the world annually. Kidney transplantation can be a lifesaving procedure and can extend a patient's life expectancy by up to 20 years. Transplant rejection is a concern for all patients though: silent or subclinical acute transplant rejection is prevalent in 25% of those who receive new kidneys within the first year. Patients may develop significant anxiety about the possibility of their organ rejecting silently, so tools for rejection management are extremely important in helping physicians secure the best outcomes for their patients.

Post kidney transplantation, the right levels of immunosuppression are key. Immunosuppressant drugs weaken a patient's immune system in order to reduce their body's reaction to the foreign organ. While the dosage may be lowered over time, patients are prescribed lifelong immunosuppressive antirejection medications, and often go through life concerned about the

stability of their transplant. **Post-transplantation patient monitoring** has typically involved both testing for immunosuppressive drug levels, which indicate whether patients are taking their medications as prescribed, as well as measuring serum creatinine levels, which indicate how well the kidney is filtering the blood. Creatinine is the current Standard of Care indicator of renal function. However, high levels of creatinine are a late indicator of damage already done to a transplanted kidney and pose unnecessary rejection risk to the patient, given that up to 50% of kidney function may already be lost before serum creatinine levels rise.

The current Standard of Care to rule out the presence of "silent" subclinical acute rejection in patients with stable renal function involves performing a "surveillance" organ biopsy. Biopsies are invasive procedures with several difficulties, from sampling error to subjective analysis, logistical challenges for patients and transplant centres, and potential complications and significant pain for the patient.

Given their high cost, they are not performed regularly. 80% return negative (i.e. normal) results, leaving patients unnecessarily subjected to the associated risks and pain of a biopsy.

Eurofins Transplant Genomics launched TruGraf® in late 2019 – the first and only blood test that can rule out "silent" subclinical acute rejection in kidney transplant recipients with stable renal function. This **non-invasive test** allows doctors to reassure patients with stable renal function that their kidney transplant is not harbouring silent rejection by assessing whether they are adequately immunosuppressed. As a result, the test avoids the need for invasive, risky and costly surveillance biopsies, and allows for early detection of rejection risk, not just when rejection is suspected.

Eurofins Transplant Genomics worked with leading researchers at Northwestern University's Comprehensive Transplant Center and The Scripps Research Institute to develop TruGraf®. Test results support

clinicians with information to optimise immune-suppressive therapy, enhance patient care and improve transplant survival.

Most recently, Eurofins Transplant Genomics launched OmniGraf™, the first and only non-invasive test panel for the earliest and most accurate view of kidney transplant rejection. OmniGraf™ combines novel genetic biomarkers by bringing together Eurofins Transplant Genomics' TruGraf® blood gene expression test and Eurofins TRAC® donor-derived cell-free DNA assays.

Eurofins TRAC® (Transplant Rejection Allograft Check) donor-derived cell-free DNA (dd-cfDNA) assay relies on non-invasive liquid biopsy to monitor the percentage of dd-cfDNA in a patient's plasma post-transplant, using Next Generation Sequencing (NGS). As donor-derived cell-free DNA is released when the transplant organ is attacked by the host's immune system, monitoring dd-cfDNA levels can indicate whether the patient is likely to be experiencing transplant rejection.

the science behind

TruGraf® is a minimally invasive biomarker test that measures differentially expressed genes in the blood of renal transplant recipients to identify patients who are likely to be adequately immunosuppressed and in doing so rule out subclinical acute rejection. TruGraf® uses RNA microarray technology to determine whether a patient has a blood gene expression profile similar to a reference population documented to have stable renal function and normal histology. Patients that do are likely adequately immunosuppressed and are classified as "TX", indicating Transplant eXcellence and patients that do not are likely to be inadequately immunosuppressed, and are classified as "not-TX".

For references, please see bibliography



FIGHTING

Limiting the spread of African Swine Fever

AN ANIMAL HEALTH ENDEMIC



ASF is caused by the **African swine fever virus (ASFV)**, which spreads among domestic pigs through contact with a sick animal, an asymptomatic carrier or a soft tick, or through the consumption of contaminated animal feed. Usually suffering a high fever and visible haemorrhages, infected pigs can also experience weight loss, pneumonia, skin ulcers, and tremors, before many fall into a comatose state and die within just a few days of infection. The impact on a pig farm is catastrophic: infection not only spreads rapidly but means that any surviving pigs in the herd must be culled to stop the spread of the virus.

In 1984, following an outbreak of ASF in Europe, Gold Standard Diagnostics Madrid, at the time Eurofins Ingenasa, developed the first commercial ELISA test capable of detecting antibodies against ASFV in serum samples of pigs and wild boar, which was evaluated according to the Office International des Epizootie (OIE) rules and validated

African swine fever (ASF) is a highly contagious and usually fatal viral disease affecting domestic pigs and wild boar. Since the first recorded description of ASF in Kenya in 1921, the virus has spread throughout Africa, Europe, the Americas and Asia, with a huge impact on the swine industry. Despite several eradication programmes, the virus has continued spreading and remains endemic in some areas. Gold Standard Diagnostics Madrid, at the time Eurofins Ingenasa, has carried out important work to fight this animal disease for more than 40 years.

by the EU Reference Laboratory. The company quickly leveraged its network of veterinary experts and deployed the **ELISA test** as part of successful national eradication programmes between 1985 and 1995. During this period, monitoring through antibody tests was conducted on all pig farms in Spain. The use of this assay, together with other diagnostic techniques and strict sanitary measures, contributed to the success of the eradication programme.

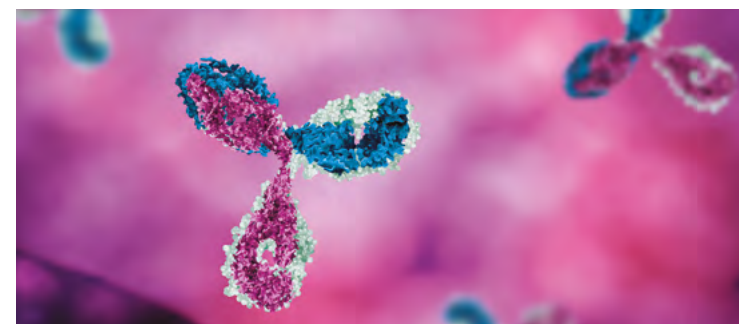
In recent decades, the virus has spread rampantly across continents again, posing a huge threat to the world's pork production. Although ASFV does not cause disease in humans, virus outbreaks are having a significant socioeconomic impact on communities that rely on the swine industry, as they result in major economic losses through the extensive culling of affected farms. For example, within seven years of ASF reaching China in 2018, millions of pigs died from the disease or had

to be culled. Stringent biosecurity management and animal quarantine measures have now become the norm in the industry, with strict limitations on animal movement and exportation during outbreaks, all taking a further toll on the economic viability of pig farming.

In response, Gold Standard Diagnostics Madrid has now developed a full catalogue of tests for specific ASFV antigen and antibody detection. The suite of tests has been used to carry out **eradication or control programmes** in for example Sardinia, the Dominican Republic, and many countries in Asia. To support improved on-site diagnosis of this disease, Gold Standard Diagnostics Madrid has enhanced its rapid tests (LFA) catalogue. The updated tests provide greater accuracy in antigen and antibody detection. Additionally, a unique in market combined test has been developed to detect both antigen

and antibody simultaneously (Combo LFA), extending the diagnostic window.

Since eradication programmes have struggled to eliminate the virus for good, control measures remain a costly necessity for pig farmers. Though no vaccine against ASFV is currently commercially available worldwide, important efforts are being carried out. Recently, Gold Standard Diagnostics Madrid participated in the European project 'A Safe DIVA (Differentiating Infected from Vaccinated Animals) Vaccine for African Swine Fever Control and Eradication (VACDIVA)', which focused on developing a safe and effective vaccine for domestic and wild swine populations, along with accompanying DIVA tests to accurately monitor their vaccination status and differentiate between infected and vaccinated animals.



the science behind

The African Swine Fever Virus is complex and variable, as a large enveloped double-stranded DNA virus, classified as the sole member of the Asfarviridae family of viruses. To test for infection with ASFV, Gold Standard Diagnostics Madrid uses different platform diagnostics, including ELISA, PCR and LFD tests.

The ELISA from Gold Standard Diagnostics Madrid is based on a monoclonal antibody, that recognises one of the most antigenic proteins of ASFV.

The INgezim® ASFV-R is an immunoenzymatic assay based on the indirect ELISA technique for the detection of specific antibodies to cp312 and p30 proteins of ASFV in serum, blood (fresh or on paper) and spleen exudate samples, from swine and wild boar.

ELISA (enzyme-linked immunosorbent assay) is a plate-based assay technique used to detect and quantify soluble substances such as antibodies. Antigens are immobilised on the plate and bind with antibodies linked to a specific enzyme. By measuring the activity of this enzyme, it can subsequently be determined whether (and how many) antibodies have bound.

Polymerase chain reaction (PCR) is a laboratory technique that detects minute traces of DNA, in this case viral DNA, in a sample. This is accomplished by multiplying the DNA present in the sample to create billions of copies, using specific enzymes and by changing temperatures.

For references, please see bibliography



Transforming cancer care through liquid biopsy

A NEW ERA OF ONCOLOGICAL DATA

For patients with cancer, regular monitoring of tumour-related genetic mutation is key to effective diagnostic, prognostic and therapeutic care. The current medical standard for molecular analysis of these mutations is tissue biopsy, an invasive and costly procedure that carries risk when used repeatedly and does not typically capture all relevant mutations in the tumour source. Eurofins Genomics and Clinical Diagnostics companies have been forerunners in sequencing tumour DNA obtained by non-invasive blood sampling and analysis, called liquid biopsy, providing specific oncological data that can have a significant impact on patient treatment decisions and outcomes.



the science behind

Personalised cancer treatments targeted at particular molecular alterations in a patient can significantly improve their prognosis and quality of life, but these therapies require that genetic monitoring of the tumour is conducted regularly. In modern medicine, this usually depends on tissue biopsy, which involves extracting suspected cancerous tissue with a small needle or by surgically removing a larger portion. However, consecutively performing this invasive procedure carries risk for the patient and is oftentimes out of the question, as well as having shortcomings in the specificity of the data it produces; due to the heterogeneity of the tissue, a single sample does not typically represent all mutations at the tumour site. For this reason, in recent years, attention in the medical community has increasingly turned to the potential of circulating tumour DNA

(ctDNA) in bodily fluids for **cancer screening and diagnostics**.

Just as healthy cells secrete DNA (called cell-free DNA, or cfDNA) into bodily fluids, such as blood, urine and cerebrospinal fluid, tumours secrete DNA in the same way. This ctDNA is characterised by genetic defects identical to and representative of the entire tumour genome. Scientists can therefore use ctDNA to genetically profile tumours with great accuracy by extracting fragments from blood plasma through a process known as **liquid biopsy** and using this to detect genetic mutations not represented in tissue samples. Though extracting DNA from plasma in a blood sample must be conducted very carefully and under specialised conditions, it is comparatively easier to do than extracting other circulating biomarkers. Liquid biopsy of ctDNA is thus a rapid, affordable and

non-invasive innovation, providing important complementary medical information on cancer progress.

In 2015, realising the great diagnostic potential of cfDNA as a biomarker, GATC Biotech, now a Eurofins Genomics company, launched the first standardised sequencing service for cfDNA, based on their experience of detecting DNA traces in blood samples for prenatal testing (see page 6: New generation of non-invasive prenatal testing methods). Today, Eurofins Genomics offers the most comprehensive **cancer gene panel** for liquid biopsy in the world, at 597 genes, focusing on major cancer types.

In 2017, Eurofins Genoma, a Eurofins Clinical Diagnostics company, launched the first commercial liquid biopsy test, Onconext™, validated and brought to market in collaboration with the Regina Elena National Cancer

Institute. Later, a Eurofins Genoma project exploring how to best manage liquid biopsy in head and neck tumours was the first of its kind to make the technique available in daily clinical practice in Italy. Currently, there is no standard for the testing method, and Eurofins Genoma was the first to prove a developed method could **detect minute traces** of cfDNA from very small, non-metastatic tumours (tumours that have not spread to other parts of the body), demonstrating the appropriateness of liquid biopsy for screening early-stage cancer, where a sensitivity at least tenfold to that of metastatic cancers is required. Today, Eurofins Genomics offers the most comprehensive cancer gene panel for liquid biopsy in the world, INVIEW Oncoprofiling at 728 genes, focusing on major cancer types.

The cfDNA sequencing services provided by Eurofins laboratories rely on either specialised Next Generation Sequencing (NGS), which can detect multiple mutations with very low allelic frequencies of less than 1%, or amplification through droplet digital PCR (ddPCR) technology. Initially, liquid biopsy entailed a complex process to distinguish ctDNA from the cfDNA secreted by healthy cells, but with the introduction of these ultrasensitive detection methods, liquid biopsy has been finetuned so that minute genetic and epigenetic aberrations can readily be detected. It's for this reason that the method favours the use of NGS and ddPCR technology over qPCR alone, the usual gold standard of diagnostics.

For references, please see bibliography



AN APPLE A DAY KEEPS THE

Detecting patulin contamination in fruit products

DOCTOR AWAY?

A study by the European Food Safety Authority points to climate change and more humid conditions as a driver of growing food safety risks, such as mycotoxin production. Although many mycotoxins are associated with grains and cereals, one of the 400 known species is particularly abundant in pome fruits like apples and pears: patulin. Eurofins Abraxis has been leading the way in the detection of patulin in commonly affected foods.

Patulin, a metabolite of the fungus *Penicillium expansum*, is produced in fruits, grains and other foods when they become infected by the fungus. Fleshy fruits, or pome fruits, and thus their derivative food products are commonly affected, putting popular products like fruit juice, apple cider, applesauce and baby foods at high risk of patulin contamination. When ingested, this mycotoxin poses major health risks to humans and animals.

As a result, more than 100 countries have adopted strict regulatory guidelines aimed at controlling the presence of patulin in food and feed products, including the US, China, and EU countries. Gold Standard Diagnostics Horsham (formerly

Eurofins Abraxis) quickly stepped up to help meet this new demand for patulin testing. In 2021, it became the first laboratory in the U.S. to develop a rapid detection method for this mycotoxin, which uses ELISA immunoassay technology to successfully detect the presence of patulin in food samples.

Additionally, Gold Standard Diagnostics Horsham is furthering its offering through the development of a lateral flow strip version of the test, to allow for even simpler qualitative or semi-quantitative results of smaller samples without the need for complex scientific equipment, another first for the industry.



the science behind

The Gold Standard Diagnostics Horsham ABRAXIS® Patulin ELISA Test Kit is suitable for the quantitative and/or qualitative detection.

ELISA (enzyme-linked immunosorbent assay) is a plate-based assay technique used to detect and quantify soluble substances. Antigens, in this case the patulin protein, are immobilised on a microtiter plate and bind with antibodies linked to a specific enzyme. By measuring the activity of this enzyme, it can subsequently be determined whether (and how many) antibodies have bound. The concentrations of the samples are determined by interpolation using the standard curve constructed with each reaction run.

The detection range of the Gold Standard Diagnostics Horsham ABRAXIS Patulin ELISA Test Kit is 7.0 ppb to 300 ppb in most fruit/juice matrices.

For references, please see bibliography

CHANGING THE GAME IN MICROBIOLOGICAL TESTING

High-throughput testing for higher quality data

Microbiological analyses are important in diverse areas, from medical research, such as antibiotic resistance research, industrial biomanufacturing, and food production in general, to the analysis of soil microbiota in farming and environmental protection efforts. However, standard microbiology testing requires intensive labour, with additional limitations on throughput and reproducibility, making it difficult to generate sufficient statistics.

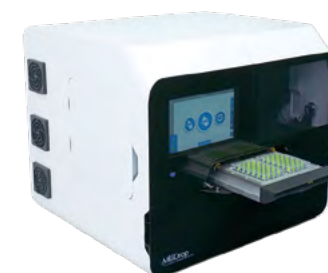
To address this, MilliDrop, now part of the Gold Standard Diagnostics network, developed a patented technology that offers an automated solution, whereby up to 2,000 biological tests can be simultaneously performed under controlled culture conditions. The innovation can also handle over 750 different conditions in a single experiment. This high-throughput solution revolutionises microbiological screening, which traditionally requires organisms (such as bacteria) to be grown on individual petri dishes.

The technology relies on the generation of droplets of 1 microlitre (one millionth of a litre) in volume, which takes place inside a capillary. The droplets are safely transported in oil, each one separated from the others by a gaseous bubble to prevent them from merging. Biological growth under the specified conditions can then be measured, with a microbial growth curve being produced for each droplet in order to obtain the growth kinetics

of the different organisms in different media. At the end of the experiment, droplets of interest can be individually recuperated for further analysis via PCR or genomic sequencing, or for regrowth.

The technology is not only high-throughput, but also excludes the risk of biological contamination during droplet preparation by preventing direct contact between the capillary tip and the solutions inside the well. Moreover, the fact that so many different conditions can be simultaneously tested in one experiment leads to reliable, reproducible data of high quality.

MilliDrop's patented technology has been tested with organisms as varied as bacteria, yeast, algae, filamentous fungi, phages and nematodes. Even before its large-scale commercial application, it has already been leveraged by multiple industrial and academic customers in Europe.



the science behind

Instead of using conventional microfluidic chips, MilliDrop has developed a cylindrical fluidic geometry where one-microlitre droplets are produced from conventional well plates, circulated inside transparent plastic capillaries, lubricated with a biocompatible oil that preserves the chemical and biological integrity of the droplets, and separated with gas bubbles (the composition of which can be adjusted) so that it can be used for specific anaerobic applications.

The continuous circulation of the droplets through a series of optical detectors (to measure biological growth by optical density or fluorescence) allows the growth kinetics indicators for each droplet to be acquired.

For references, please see bibliography



HOW SAFE IS OUR DRINKING WATER? WATER?

Detecting an emerging disinfection by-product in drinking water



Ensuring that public water systems provide clean and safe drinking water is a fundamental part of safeguarding health. To prevent the spread of waterborne diseases, water supplies are treated with disinfectants, but these chemicals themselves can form by-products that may be harmful at higher concentrations. Chloronitramide anion (Cl-N-NO_2^-) is one of the most recently identified emerging disinfection by-products (DBP) identified in drinking water treated with chloramines. In the two months following this discovery, US-based Eurofins Environment Testing Eaton Analytical developed a cutting-edge testing method to detect chloronitramide anion.

For over 100 years, chlorination has been the primary way of inactivating pathogens in water to prevent waterborne diseases. However, chlorine reacts with organic matter, bromide and iodide in water, leading to the creation of DBPs that are associated with known health risks; authorities in many countries have therefore introduced regulations concerning chlorination. As a result, more and more public water systems are moving towards using alternative disinfectants, such as inorganic chloramines, to avoid DBP contamination. Approximately one-third of water utilities in the US currently use chloramination as their primary or secondary means of disinfection of drinking water. In late 2024, new research has found that inorganic chloramines also form their own by-products, with a potential impact on human health.

One of these by-products is chloronitramide anion, which was unidentified by scientists until late 2024, when a groundbreaking study published in *Science*, by Fairey et al. highlighted the presence of chloronitramide anion in chloraminated drinking water at a concentration of up to 120 µg/L. Although its toxicity is still being investigated, the fact that chloronitramide anion contains chlorine and a nitrogen-nitrogen bond, and shares structural similarities with some highly toxic compounds, raises important questions about the potential threat it poses to human health.

Within two months of the study's publication, Eurofins Environment Testing Eaton Analytical partnered with its author, Dr. Julian Fairey from the University of Arkansas, to develop a method to analyse chloronitramide

anion in drinking water. This method leverages state-of-the-art technology for enhanced accuracy and provides ultra-sensitive detection down to 0.1 µg/L. Using this new method, chloronitramide anion was detected in all chloraminated drinking water samples tested, at a concentration of up to 200 µg/L. In addition, almost all chlorinated drinking water samples were found to contain chloronitramide anion at low µg/L concentrations.

This new method will provide water utilities with important data on chloronitramide anion contamination, ahead of potential regulatory changes.



the science behind

Eurofins Environment Testing Eaton Analytical used the "reversed-phase (RP) LC/MS/MS" testing methodology, which presents several advantages over the more commonly used ion chromatography (IC) or so-called hydrophilic interaction liquid chromatography tandem mass spectrometry (HILIC/MS/MS) technology – for instance, issues with specificity, resolution, and matrix interferences. Eurofins Environment Testing Eaton Analytical's method employs ^{15}N -labelled analyte natives as the internal standard, as it responds to the analysis in the exact same way as the target analyte. Therefore, this internal standard technique can compensate for matrix interferences and analytical variations and help achieve optimal sensitivity, accuracy, and precision in complex real-world drinking water samples that vary in hardness, alkalinity, and solids. In addition to monitoring the masses of chloronitramide anion for quantitation, extra masses specific to chloronitramide anion were also monitored to confirm sample results, which further ensured the accuracy of the data.

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