Prostate cancer (PCa) is the leading cause of cancer-related deaths in males. Patients with PCa can go undiagnosed for many years as the symptoms mostly appear when the tumour has grown large enough to put pressure on the urethra. Currently, prostate-specific antigen (PSA) is used as a marker to help detect and predict PCa onset. However, PSA is not specific to PCa, and PSA is also elevated in other prostatic conditions. This lack of specificity in PCa screening can lead to over-diagnosis, invasive biopsies, and anxiety in patients.

This highlights the need for improving PCa screening to get an accurate diagnosis and avoid unnecessary procedures. This year’s EAU21 Congress invited experts involved in some of the latest state-of-the-art PCa screening trials to discuss their results and the implications for future screening strategies. Six speakers shared their preliminary and ongoing data from clinical trials and their methods for implementing a more efficient PCa screening process.

The first speaker, Monique J. Roobol, Erasmus University Medical Centre, Cancer Institute, Rotterdam, the Netherlands, gave an informative introduction on the burden of PCa screening. Roobol’s team developed a PCa screening algorithm that assessed the risk of PSA screening in individuals. The algorithm takes into consideration the geography of the patient and individual risk factors such as age. The algorithm showed that only 10% of males aged 50–59 years and only 25% of males aged 60–70 years would move on to risk stratification.

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The results showed that current methods of PCa testing are not suitable for all individuals, and a tailored approach is urgently needed. Roobol also added that stopping testing in elderly individuals with low PSA levels would reduce unnecessary harm and costs. Finally, Roobol shared the vision of the European Cancer Organisation (ECCO) and explained that they aim to provide all males with access to risk-based screening programmes in Europe by 2027.

The next speaker was Christian Arsov, Department of Urology, University Hospital Dusseldorf, Germany, who discussed the preliminary results from the ongoing PROBASE trial. The randomised study took place in four regions of Germany and involved 46,642 males aged 45 years old. There were two study arms: study arm A had an immediate screening of baseline PSA at age 45 years, and study arm B deferred screening until the age of 50 years. Overall, 344 men were in the high-risk group as they had PSA levels above 3. The participants underwent a second PSA test to confirm the PSA levels, and 179 men were still at risk. The results of the trial showed a very low prevalence of PCa in 45-year-olds (0.2%). The speaker emphasised that elevated PSA levels should always be confirmed with a second test.

Over the years, PCa screening has evolved, and the screening steps are as follows: PSA test, systemic biopsy, MRI, and finally, targeted biopsy. At each stage, a percentage of individuals can be ruled out, therefore reducing the chance of over-diagnosis.

Anssi Auvinen from Tampere University, Finland, discussed the ongoing PROSCREEN trial. In this trial, the researchers wanted to add another step to PCa screening called the 4K score, a combination of four blood tests, after the PSA test to further eliminate males who are not eligible for an MRI or biopsy. The 4K score eliminated 30% of participants. This trial is still in its infancy and experienced delays due to the pandemic. Calculations show that in 15 years, the researchers should be able to establish the efficacy of adding the 4K score to PCa screening.

The next speaker, Tobias Nordstrom, Karolinska Institute, Stockholm, Sweden, also aimed to improve the risk-adapted screening process by combining a genomic test with MRI-targeted
biopsy in the STHLM3 MR Phase II trial. Stockholm-3 (STHLM3) is a blood-based test that involves a combined assessment of PSA, four proteins, 101 genetic markers, and medical history, i.e., earlier biopsies. All 12,750 respondents who took part in this trial had a PSA and STHLM3 test. Participants were randomised to receive either a traditional biopsy or MRI. Interestingly, the results showed that STHLM3 discriminated better than PSA for significant cancer, with an area under the curve of 0.76 compared to 0.60 in PSA. Nordstrom concluded that combining STHLM3 with MRI targeted biopsy reduced over-detection yet still effectively detected significant cancer.

This was followed by a presentation by Jacob Fredsoe, Aarhus University Hospital, Denmark. Fredsoe explained the PROCARIS trial, which took a unique approach using genetic markers, specifically single-nucleotide polymorphisms (SNP), to assess PCa risk. To reduce the likelihood of over-diagnosis, Fredsoe’s group explored the option of reducing repeated PSA tests depending on an individual’s genetic risk.

This study was conducted in central Denmark and involved 146 general practices; n=73 general practices were intervention practices, and n=73 were control practices. Men who visited their general practitioners were given a PSA test and offered a genetic test. If the genetic test showed they were at normal risk, they had no further PSA tests. However, if the genetic test showed they were at high risk, the patient was encouraged to have yearly PSA tests.

The primary endpoint was the proportion of males with repeated PSA tests within 2 years. The results showed that males with normal genetic risk had a lower chance of repeated PSA tests, and males with high genetic risk were more likely to have elevated PSA, biopsy, and a PCa diagnosis.

The final speaker, Robert Nam, Sunnybrook Research Institute, Toronto, Canada, discussed the ‘Risk-Adapted Screening with MRI Only: MVP Trial.’ In PCa screening, MRI is often used alongside PSA tests or genomic tests. However, Nam’s group aimed to explore whether MRI could be used as a stand-alone tool to detect PCa. In this Phase III study, patients aged 50 years or older underwent either prostate MRI or PSA test in a randomised 1:1 ratio.

Collectively, 525 were able to participate in this trial: 246 had an MRI, and 248 had a PSA test. Surprisingly, 18 patients dropped out of the study as soon as they found out they would be getting a PSA test as they were upset and were hoping to be in the MRI group. Preliminary results showed that the MRI arm of the study detected more than double the rate of PCa compared to the PSA arm. Nam concluded that MRI is a lot more selective than PSA tests and avoids over-treatment and over-diagnosis. Regardless of initial expenses, it would work out to be more cost-effective in the long term to have an MRI scan rather than repeated PSA tests.

Despite a variety of clinical trials in PCa screening discussed in this year’s congress, there was a common theme: a risk-adapted strategy to screening. The speakers proposed novel approaches to get the best out of the screening process whilst reducing harm and over-diagnosis. Many of these trials were in the beginning stages and only showed preliminary results; however, the speakers agreed this was a step in the right direction. In the future, other unanswered questions may be explored such as cost, convenience, and anxiety. Some of the results from these trials will not be confirmed for another decade, but the researchers are optimistic that there will be exciting improvements in PCa screening in the near future.