

# Translation of DNA methylation markers for the early detection of renal cell cancer

Citation for published version (APA):

Lommen, K. W. J. M. P. (2022). Translation of DNA methylation markers for the early detection of renal cell cancer: raising the odds. [Doctoral Thesis, Maastricht University]. Maastricht University. https://doi.org/10.26481/dis.20221017kl

Document status and date: Published: 01/01/2022

DOI: 10.26481/dis.20221017kl

**Document Version:** Publisher's PDF, also known as Version of record

## Please check the document version of this publication:

 A submitted manuscript is the version of the article upon submission and before peer-review. There can be important differences between the submitted version and the official published version of record. People interested in the research are advised to contact the author for the final version of the publication, or visit the DOI to the publisher's website.

• The final author version and the galley proof are versions of the publication after peer review.

 The final published version features the final layout of the paper including the volume, issue and page numbers.

Link to publication

### General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these riahts.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
  You may freely distribute the URL identifying the publication in the public portal.

If the publication is distributed under the terms of Article 25fa of the Dutch Copyright Act, indicated by the "Taverne" license above, please follow below link for the End User Agreement:

www.umlib.nl/taverne-license

#### Take down policy

If you believe that this document breaches copyright please contact us at:

repository@maastrichtuniversity.nl

providing details and we will investigate your claim.

## **IMPACT PARAGRAPH**

In 2020, approximately 430.000 new cases of kidney cancer were reported globally, representing 2.2% of all cancers diagnosed<sup>1</sup>. In that same year, an estimated 180.000 deaths could be attributed to kidney cancer<sup>1,2</sup>. Renal cell carcinoma (RCC) is the most prevalent type of kidney cancer, responsible for 90-95% of all cases, and the incidence rates have been rising with 2% per year over the past two decades<sup>2,3</sup>. The 5-year survival rate of RCC confined to the kidney is 93%, whereas the 5-year survival of distant metastasized RCC is decreased to 13%<sup>4</sup>. Globally, 3.3 million disability-adjusted life-years were estimated for RCC in 2017, and the disease burden has not decreased over the past 30 years despite advancements in RCC management<sup>5,6</sup>. With the increasing use of imaging techniques, a substantial proportion of patients are diagnosed after a coincidental finding during an unrelated procedure<sup>7,8</sup>. The rising incidence, partly caused by the increasing amount of incidental findings, together with the aging population will continue to increase both disease and economic burden because of RCC.

The increasing amount of incidentally diagnosed small renal masses (SRMs), and the fact that only 50-70% of these SRMs are correctly being diagnosed as benign or malignant based on medical imaging, goes hand-in-hand with overtreatment of many patients<sup>2,4,7,8</sup>. Whenever the nature of an SRM is uncertain, partial or radical nephrectomy can be decided upon. Patients will undergo burdensome surgeries to remove (part of) a kidney, never without risks of complications, which after pathological evaluation turn out to have been unnecessary in 25% of cases<sup>9</sup>. In addition, patients experience current diagnostic procedures like computed tomography (CT) scans and magnetic resonance imaging (MRI) scans as unpleasant because of scan duration, loud noise and space- and motion restrictions<sup>10,11</sup>. Next to that, these imaging procedures are considered timeconsuming, not only because of the duration of the scan itself, but also because patients have to travel to and from the hospital<sup>10</sup>. From a hospital- and societal burden perspective, imaging techniques are also burdensome in terms of costs, time, effort and the need for specialized staff. Taken together, this emphasizes the importance of improving the current regime in accurately diagnosing RCC in early stages. To limit and potentially substitute part of burdening diagnostic imaging procedures, more accurately diagnose SRMs and avoid overtreatment, studies have been aiming to improve RCC diagnostics by focusing on non-invasive molecular markers in liquid biopsies. These patient-friendly diagnostic biomarkers for RCC could reduce both health- and economic burden not only by directly replacing costly and unpleasant imaging procedures, but also by avoiding unnecessary surgeries and subsequent medical follow-up.

Despite the great interest and research invested in molecular markers to replace invasive procedures, only <1% of all published biomarkers have reached clinical care<sup>12,13</sup>. This is in line with the fact that a large proportion of research investments being avoidably wasted at this moment<sup>14</sup>. Unfortunately, the same holds true for DNA methylation biomarkers specifically, illustrated by the fact that not a single DNA methylation biomarker for diagnosing RCC has reached clinical care.

After initial publication, the vast majority of biomarkers never reach the stage of validation. In this thesis, we have addressed several reasons for the lack of success and validation and illustrated the effect of assay type, assay design, sample type, sample selection and control selection on biomarker development in the individual chapters of this thesis. In addition, we discussed to adequately address all these problems when performing a biomarker study. As argued in **Chapter 7**, validation studies are considered less innovative as compared to identification of novel biomarkers, and therefore also less of interest to scientific journals and funding agencies, leading to publication bias and an imbalance in funding streams<sup>15</sup>. Changing this scientific mentality is a shared responsibility of the academic community, including researchers, peer reviewers, journals, editors, universities and funding organizations. Filling knowledge gaps, rather than creating even more of them needs to be supported, facilitated and enforced by all parties. Accordingly, reducing research waste starts with many considerations upfront, rather than merely in the lab.

Suggestions provided in this thesis strive towards more efficient use of biomarker research funding, thereby not only reducing research waste, but also increasing the chance of developing clinically useful biomarkers for the accurate and early diagnosis of RCC. In that way, the results and perspectives from this thesis have both scientific and economic impact, as changing the biomarker research mentality will encourage decent and reproducible research, thereby facilitating validation of biomarkers, and subsequently allowing clinical utility of these biomarkers. The clinical applicability of these biomarkers will in turn lead to a reduction in health- and economic burden, by replacing and sparing imaging procedures, surgeries and medical follow-up.

## REFERENCES

- Sung H, Ferlay J, Siegel RL, et al. Global Cancer Statistics 2020: GLOBOCAN Estimates of Incidence and Mortality Worldwide for 36 Cancers in 185 Countries. *CA: A Cancer Journal for Clinicians*. 2021; 71(3):209-249.
- Du Z, Chen W, Xia Q, Shi O, Chen Q. Trends and projections of kidney cancer incidence at the global and national levels, 1990–2030: a Bayesian age-period-cohort modeling study. *Biomarker Research*. 2020;8(1):16.
- **3.** Woo S, Cho JY. Imaging findings of common benign renal tumors in the era of small renal masses: differential diagnosis from small renal cell carcinoma: current status and future perspectives. *Korean J Radiol.* 2015;16(1):99-113.
- Society AC. Cancer facts & figures 2021. 2021. https://www.cancer.org/content/dam/cancer-org/ research/cancer-facts-and-statistics/annual-cancer-facts-and-figures/2021/cancer-facts-andfigures-2021.pdf.
- 5. Safiri S, Kolahi A-A, Mansournia MA, et al. The burden of kidney cancer and its attributable risk factors in 195 countries and territories, 1990–2017. *Scientific Reports*. 2020;10(1):13862.
- 6. Zi H, He S-H, Leng X-Y, et al. Global, regional, and national burden of kidney, bladder, and prostate cancers and their attributable risk factors, 1990–2019. *Military Medical Research*. 2021;8(1):60.
- 7. Battagli C, Uzzo RG, Dulaimi E, et al. Promoter hypermethylation of tumor suppressor genes in urine from kidney cancer patients. *Cancer research*. 2003;63(24):8695-8699.
- Rydzanicz M, Wrzesinski T, Bluyssen HA, Wesoly J. Genomics and epigenomics of clear cell renal cell carcinoma: recent developments and potential applications. *Cancer letters*. 2013;341(2):111-126.
- 9. Ambani SN, Wolf JS, Jr. Renal mass biopsy for the small renal mass. Urol Oncol. 2018;36(1):4-7.
- **10.** Oztek MA, Brunnquell CL, Hoff MN, et al. Practical Considerations for Radiologists in Implementing a Patient-friendly MRI Experience. *Topics in Magnetic Resonance Imaging*. 2020;29(4).
- 11. Rosenkrantz AB, Pysarenko K. The Patient Experience in Radiology: Observations From Over 3,500 Patient Feedback Reports in a Single Institution. *J Am Coll Radiol*. 2016;13(11):1371-1377.
- 12. Poste G. Bring on the biomarkers. *Nature*. 2011;469(7329):156-157.
- 13. Kern SE. Why your new cancer biomarker may never work: recurrent patterns and remarkable diversity in biomarker failures. *Cancer research*. 2012;72(23):6097-6101.
- 14. Macleod MR, Michie S, Roberts I, et al. Biomedical research: increasing value, reducing waste. *Lancet*. 2014;383(9912):101-104.
- Ioannidis JPA, Bossuyt PMM. Waste, Leaks, and Failures in the Biomarker Pipeline. *Clin Chem.* 2017; 63(5):963-972.