A scoping review and meta-analysis on the prevalence of pan-tumour biomarkers (dMMR, MSI, high TMB) in different solid tumours

*Appendix II. Managing overlap of data sources for meta-analyses and studies

*with overlapping data sources*

**RUNNING TITLE:** Prevalence of dMMR, MSI and high TMB in different solid cancers

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1. Managing overlap of data sources for meta-analyses

a) Some patient cohorts have been included in biomarker prevalence estimates in multiple original research studies or systematic reviews. To avoid data duplication in the meta-analyses in this review, we identified studies with overlapping data sources and only included at most one estimate based on a specific patient cohort.

b) For each biomarker and each major data source that was included multiple original research studies on the prevalence of dMMR/MSI/high TMB in specific cancer(s) and pan-cancer analyses (e.g., Foundation Medicine database, Memorial Sloan Kettering Cancer Centre patients, analysis of the Cancer Genome Atlas), we used the following approach.

- Pan-cancer overall prevalence: the study with the largest sample size was included in the meta-analysis.
- Cancer-specific overall prevalence: the study with the largest sample size for the specific cancer was included in the meta-analysis for “overall” cancer-specific estimates.
- Cancer-specific prevalence for early-stage or advanced-stage cancers: the study with the largest sample size for the specific cancer type and stage was included in the meta-analysis.

c) Systematic reviews and/or meta-analyses

- For some cancers (e.g., colorectal cancer, endometrial cancer), we identified multiple systematic reviews and/or meta-analyses that reported the prevalence of dMMR/MSI/high TMB to address a specific research question. For each cancer type, we included the study with the largest sample size. If the sample size was similar across multiple systematic reviews and/or meta-analyses, we included the study for which the research question was most aligned with our scoping review.
- To avoid further data duplication, if a meta-analysis in this review included estimates from a previously published meta-analysis, the underlying original studies included in the previously published meta-analysis were excluded from the corresponding meta-analysis in this review.

See Section 2 below for the list of studies with overlapping data sources that reported the prevalence of dMMR/MSI/high TMB and the rationale for their inclusion or exclusion in meta-analyses.
2. Studies with overlapping data sources and the rationale for their inclusion or exclusion in meta-analyses

1) Studies reporting the prevalence of dMMR/MSI/high TMB based on data from the Cancer Genome Atlas

| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|---------------|-----------|---------------------|-----------------|---------------------------------------|------------------------|----------------------------------|
| Fan (2020)¹   | Gastric cancer | Overall            | • Period N/S    |
|               |           |                     | • Total (N=924) |
|               |           |                     | o TCGA (n=440)  |
|               |           |                     | o Local hospitals (n=484) | TMB | MSI | • The prevalence of high TMB was not included in the data synthesis due to a data-driven high TMB cut-off (75th percentile) |
|               |           |                     | • The combined prevalence of MSI from both the TCGA and two local hospitals were included in the data synthesis |
|               |           |                     | o The TCGA cohort is larger than that included in Qu et al.² and Ren et al.³ | |
| Qu et al. (2020)² | Gastric cancer | Overall            | • Period N/S    |
|               |           |                     | • Total (N=386)  |
|               |           |                     | • MSI | Excluded from the data synthesis due to the smaller sample size than Fan et al.¹ |
| Ren et al. (2020)³ | Gastric cancer | Overall            | • Period N/S    |
|               |           |                     | • Total (N=383)  |
|               |           |                     | • TMB | MSI | • Prevalence of TMB was provided in graphical format only and excluded from the data synthesis |
|               |           |                     | • Prevalence of MSI was excluded from the data synthesis due to the smaller sample size than Fan et al.¹ |
| Li (2020)⁴    | Gastric cancer | Overall            | • Period N/S    |
|               |           |                     | • Total (N=510)  |
|               |           |                     | o TCGA (n=210)  |
|               |           |                     | o GEO (n=300)   | MSI | Excluded from the data synthesis |
|               |           |                     | o The TCGA cohort is likely to be a subset of the cohort reported in Fan et al.¹ |
|               |           |                     | o The GEO cohort did not satisfy the minimum cancer-specific sample size cut-off (400+) |
| Dai (2020)⁵   | Gastric cancer | IB-III              | • Period N/S    |
|               |           |                     | • Total (N=424)  |
|               |           |                     | o TCGA (n=202)  |
|               |           |                     | o ACRG (n=138)  |
|               |           |                     | o Local (n=89)  | MSI | Included in the data synthesis for early-stage gastric cancers only |

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; TCGA = The Cancer Genome Atlas; ACGR = Asian Cancer Research Group; GEO = Gene Expression Omnibus.

¹ Number of samples available for the biomarker status.
2) Studies reporting the prevalence of dMMR/MSI/high TMB based on data from the Foundation Medicine Database

| Author (year)     | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion                                                                                                                                 |
|-------------------|-----------|---------------------|-----------------|---------------------------------------|------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Chan (2019)       | Pan-cancer| 30 cancer types     | Overall         | • Period N/S                           | TMB (≥10 mut/Mb)       | • Excluded from the data synthesis since the prevalence of high TMB was provided in graphical format only                                                                                     |
| Trabucco (2019)   | Pan-cancer| 34 solid tumours    | Overall         | • Period N/S                           | MSI                    | • Excluded from the data synthesis                                                                                                      |
|                   |           | ~10 haematopoietic  |                 | • Total (N=67,644)                     |                        | o Cancer-specific prevalence was shown only for cancer types with ≥100 samples, all of which had smaller sample size than Yoshino et al.                                                                 |
|                   |           | tumours             |                 |                                        |                        |                                                                                                                                                                                                  |
| Yoshino (2020)    | Pan-cancer| 30 adult tumours    | Overall         | • Period N/S                           | MSI                    | • Cancer-specific and pan-cancer prevalence of both MSI and of high TMB (≥20 mut/Mb) were included in the data synthesis due to the largest sample size, after excluding haematologic tumours and lymphoma |
|                   |           | 10 Paediatric       |                 | • Adult (N=164,410)                    | TMB (≥20 mut/Mb)       |                                                                                                                                                                                                  |
|                   |           | tumours             |                 | • Paediatric (N=3,592)                 |                        |                                                                                                                                                                                                  |
| Huang (2021)      | Pan-cancer| 6 tumour groups     | Overall         | Jan. 2016 - Nov. 2019                  | MSI                    | • Prevalence of MSI was excluded from the data synthesis due to the smaller sample size than Yoshino et al.                                                                                     |
|                   |           | encompassing multiple common cancer types |     | • Total (N=48,782)                     | TMB (≥10 mut/Mb)       | • Prevalence of high TMB (≥10 mut/Mb) was included in the data synthesis                                                                                                                     |
|                   |           | 9 cancer types      |                 |                                        | TMB (≥20 mut/Mb)       | o Cancer-specific prevalence: soft tissue sarcoma, melanoma, head and neck cancer, NSCLC, bladder/urothelial cancer, breast cancer, cervical cancer, endocrine tumour and neuroendocrine tumour (Note: Endocrine tumours were included in cancer-specific analysis due to thyroid cancer being the major cancer type. Neuroendocrine tumours are rare and were included in cancer-specific analysis) |
|                   |           |                     |                 |                                        |                        | o Pan-cancer prevalence                                                                                                                                                                         |
|                   |           |                     |                 |                                        |                        | o Tumour group-specific prevalence: CNS tumours, gastrointestinal cancers, genitourinary tract cancers, gynaecological cancers, excluding esophageal SCC (rare histologic sub-type) and cancers not otherwise specified |
|                   |           |                     |                 |                                        |                        | • Prevalence of high TMB (≥20 mut/Mb) was excluded from the data synthesis due to the smaller sample size than Yoshino et al.                                                             |
| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size<sup>a</sup> | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|---------------------|----------------|-----------------------------------|------------------------|--------------------------------|
| Goodman (2019)<sup>10</sup> | Multiple cancer types | SCC only | Overall | Period N/S | MSI | • Specimens from metastatic site and primary tumour site represented 35.9% and 46.1%, respectively, with NSCLC representing 34.2% of the cohorts |
| Parikh (2019)<sup>11</sup> | Gastrointestinal cancers | Tubular only | Advanced | Period N/S | TMB (≥20 mut/Mb) | • Excluded from the data synthesis since this study focused on specific histologic sub-type only |
| Necchi (2020)<sup>12</sup> | Bladder/urothelial cancer | Urothelial carcinoma | Advanced | June 2012 – July 2018 | MSI | • Included in the data synthesis since this study reported the prevalence of both MSI and high TMB (≥10 mut/Mb, ≥20 mut/Mb) in advanced bladder/urothelial cancers |
| Necchi (2020)<sup>13</sup> | Bladder/urothelial cancer | Urothelial carcinoma | Advanced | Aug. 2014 – Nov. 2018 | MSI | • Excluded from the data synthesis since Necchi et al.<sup>12</sup> reported the prevalence of both MSI and high TMB (≥10 mut/Mb, ≥20 mut/Mb) in advanced bladder/urothelial cancers |
| Madison (2020)<sup>14</sup> | Bladder/urothelial cancer | Urothelial carcinoma | Advanced | Period N/S | MSI | • Excluded from the data synthesis since Necchi et al.<sup>12</sup> reported the prevalence of both MSI and high TMB (≥10 mut/Mb, ≥20 mut/Mb) in advanced bladder/urothelial cancers |
| Chung (2019)<sup>15</sup> | Prostate cancer | Overall | Period N/S | Total (N=3,476) | MSI | • Prevalence of both MSI and high TMB (≥20 mut/Mb) was excluded in the data synthesis due to the smaller sample size than Yoshino et al.<sup>8</sup> |
| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|---------------------|----------------|---------------------------------------|-------------------------|----------------------------------|
| Necchi (2020) | Testicular cancer | Germ cell tumours only | Advanced (relapsed after CT) | 2012 - 2017 Total (N=107) o Seminoma (n=23) o Non-seminoma (n=84) | MSI TMB (≥10 mut/Mb) TMB (≥20 mut/Mb) | Prevalence of high TMB (≥10 mut/Mb) was included in the data synthesis after combining estimates from both primary and metastatic site given metastatic sites include lymph node metastasis only and not limited to distant metastasis o Primary site (3.9%), metastatic site (6.2%) |
| Patel (2020) | Brain tumour | Paediatric tumours only (age ≤ 21 years) | Overall | Nov. 2012 – May 2017 Total (N=723) o HGG (n=277) o LGG (n=235) o MB (n=134) o Others (n=77) | TMB (≥20 mut/Mb) | Excluded from the data synthesis o Likely to be a subset of the cohort reported by Yoshino et al.8 (408 non-gliomas and 800 gliomas in paediatric patients) o Yoshino et al.8 was the only one study reported the prevalence of dMMR/MSI/high TMB and data synthesis in the prevalence of the pan-tumour biomarkers in paediatric solid tumours was not performed |
| Chow (2020) | Sarcoma | Soft tissue sarcoma o DSRCT | Overall | 2012-2018 Total (N=83) | MSI TMB (≥20 mut/Mb) | Excluded from the data synthesis o Focused on a rare histologic sub-type o Yoshino et al. (2020)8 reported the prevalence of TMB (≥20 mut/Mb) in soft tissue sarcoma |
| Eskander (2020) | Lung cancer o Cervical cancer | Lung: SCLC o Cervix: HGNECC | Overall | Mar. 2013 – Dec. 2017 Total (N=1,800) o SCLC (n=1800) o HGNECC (n=97) | MSI TMB (≥20 mut/Mb) | Excluded from the data synthesis o SCLC: a sub-set of Foundation Medicine Database cohort reported by Yoshino et al. (2020)8 o HGNECC: A rare histologic sub-type of cervical cancer |
| Author (year)       | Cancer(s)                                      | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|---------------------|------------------------------------------------|---------------------|-----------------|---------------------------------------|-------------------------|----------------------------------|
| Singhi (2019)       | Pancreatic cancer                              | Pancreatic ductal adenocarcinoma | Overall         | Period N/S Total (N=3,594)            | MSI, TMB ≥20 mut/Mb      | Prevalence of MSI was excluded from the data synthesis since this study was included in a systematic review by Luchini et al.21. Prevalence of TMB (≥20 mut/Mb) was included in the data synthesis given Yoshino et al.8 did not report the prevalence of TMB (≥20 mut/Mb) in pancreatic cancer. |
| Huang (2020)        | Breast cancer                                  | HR+/HER2-, HER2-, TNBC   | Overall         | Mar. 2019s – June 2019 Total (N=312) | TMB ≥9 mut/Mb           | Exclude the data synthesis due to the uncommon high TMB cut-off. |
| Sivapiragasam (2021) | Breast cancer                                  | ER+/HER2-, ER-/HER2+, TNBC | Metastatic      | Sep. 2012 – July 2018 Total (N=3,831) | MSI, TMB (≥10 mut/Mb) TMB (≥20 mut/Mb) | Included in the data synthesis given this study focused on metastatic breast cancer only, after combining the estimates of any molecular subtypes o MSI-H: ER+/HER2- (2/1237, 0.2%), ER-/HER2/amp (2/1953, 0.1%), TNBC (3/641, 0.5%); reported % is 0.4% o TMB≥10 mut/Mb: ER+/HER2- (99/1237, 8%), ER-/HER2/amp (234/1953, 12%), TNBC (58/641, 9%) o TMB≥20 mut/Mb: ER+/HER2- (25/1237, 2%), ER-/HER2/amp (39/1953, 2%), TNBC (19/641, 3%) |
| Ross (2020)         | Cancer of unknown primary                      | Overall              | Period N/S Total (N=303) | MSI, TMB (≥16 mut/Mb) | Excluded from the data synthesis due to the smaller sample size than Yoshino et al.8 |
| Shao (2020)         | Multiple cancer types (same 10 rare solid tumour types included in KEYNOTE 158 study) | Lung (SCLC, mesothelioma), Cervical cancer, Anal cancer, Vulvar cancer, Endometrial cancer, Biliary tract cancer, Thyroid cancer, Salivary gland carcinoma | Overall | ~July 2018 Total (N=2,992) | TMB (≥10 mut/Mb) | Excluded from the data synthesis o The cohort was generated by linking the Flatiron Health electronic health records database to the Foundation Medicine database of tumour sequencing results (Flatiron Health-Foundation Medicine Clinicogenomic Database), which is likely to be a subset of the study cohort reported by Huang et al.2. |
| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample sizea | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|---------------------|-----------------|----------------------------------------|------------------------|---------------------------------|
| Singal (2019) | Lung cancer | NSCLC | Advanced | • Jan. 2011 – Jan. 2018  
• Total (N=4,064) | • TMB (≥20 mut/Mb) | Included in the data synthesis since the study focused on advanced NSCLC only, although the study cohort was identified from the Flatiron Health-Foundation Medicine Clinicogenomic Database |
| Okamura (2020) | 9 cancer types associated with high ARID1A alterations (>5%) | Lung cancer  
Colorectal cancer  
Breast cancer  
Melanoma  
Pancreatic cancer  
Cholangiocarcinoma/hepatocellular  
Gastric/esophageal cancer  
Endometrial cancer  
Urothelial bladder carcinomas | Overall | • Period N/S  
• Total (N=1,093) | • MSI  
• TMB (≥20 mut/Mb) | Excluded from the data synthesis  
• Tissue DNA from the UCSD was analysed by Foundation Medicine, which is likely to be a sub-set of the study cohort reported by Yoshino et al.8  
• Even if this is not a sub-set of the Foundation Medicine cohort, inclusion of this study will not make substantial difference due to small cancer-specific sample size. For example, sample size for colorectal, endometrial, bladder and gastric/esophageal cancer are below the minimum sample size cut-off. |

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; CNS = central nervous system; SCC = squamous cell carcinoma; ADC = adenocarcinoma; NSCLC = non-small cell lung cancer; UCSD = University of California San Diego; FM = Foundation Medicine; HGG = high grade glioma; LGG = low grade glioma; MB = medulloblastoma; CT = chemotherapy; DSRCT = desmoplastic small round cell tumour; SCLC = small cell lung cancer; HGNECC = high grade neuroendocrine cervical cancer; HR+ = hormone receptor positive; HER2- = human epidermal growth factor receptor negative; TNBC = triple negative breast cancer; UCSD = University of California San Diego; CRC = colorectal cancer.

a Number of samples available for the biomarker status.
### 3) Studies reporting the prevalence of dMMR/MSI/high TMB based on data from the Memorial Sloan Kettering Cancer Center

| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|---------------|-----------|---------------------|----------------|---------------------------------------|------------------------|----------------------------------|
| Latham (2019) | Pan-cancer | 50+ cancer types    | Overall        | Jan. 2014 – June 2017, Total (N=15,045) | MSI                    | Excluded from the data synthesis  |
|               |           |                     |                |                                       |                        | o Pan-cancer prevalence: Hechtman et al. reported the similar pan-cancer prevalence of MSI with the larger sample size (2.2% [326/15045] vs 2.0% [582/29530]) |
|               |           |                     |                |                                       |                        | o Cancer-specific prevalence: Prevalence of high-frequency (MSI-H) or indeterminate microsatellite instability (MSI-L) was reported. Prevalence of MSI-H and MSI-L was separately presented in graphical format only |
|               |           |                     |                |                                       |                        | • Breast (n=2,371) and lung (n=1,952) cancers represent 28.7% of tumours and CRC and EC represent 9% of all tumours (n=1,351) |
| Hechtman (2020) | Pan-cancer | Overall        | 2014 - 2018, Total (N=29,530) | MSI                    | Pan-cancer prevalence of MSI only was included in the data synthesis due to the largest sample size |
|               |           |                     |                |                                       |                        | o Cancer-specific prevalence was not provided since this study focused on a sub-group whose IHC results for dMMR was available. Discordance between MSI and dMMR was 7.2% (overall), 6.4% (CRC and 4.9% (EC) |
| Valero (2021) | Pan-cancer | 17 cancer types   | Overall        | Period N/S, Total (N=10,233) | MSI, TMB (percentiles) | Excluded from the data synthesis  |
|               |           |                     |                |                                       |                        | o Prevalence of MSI: Pan-cancer prevalence (3%, 264/10233) was provided with the smaller sample size than reported by Hechtman et al. |
|               |           |                     |                |                                       |                        | o Prevalence of high TMB: data driven high TMB cut-off (percentiles) was used |
| Jimenez-Rodriguez (2021) | Colon cancer | Adenocarcinoma | I/II/III | Feb. 2007 – Dec. 2014, Total (N=443) | dMMR | Included in the data synthesis since this study focused on early-stage colon cancer only |
| Middah (2019) | CRC       | Overall        | Jan. 2014 – Oct. 2017, Total (N=1,751) | MSI | Included in the data synthesis |
| Greally (2019) | Esophagogastric cancer | • Esophageal/GE junction cancer, • Gastric cancer | Metastatic | Sep. 2013 – May 2018, Total (N=161) | dMMR/MSI | Excluded from the data synthesis  |
|               |           |                     |                | o Esophageal/GE junction (n=85) |                        | o Prevalence of either dMMR or MSI measured by IHC or selected gene panel testing was provided without specifying denominators |
| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|---------------------|-----------------|---------------------------------------|------------------------|---------------------------------|
| Audenet (2019)34 | Bladder/urothelial cancer | Urothelial carcinoma only | Overall | Period N/S  
Total (N=649)  
o Bladder (n=454)  
o Upper urinary tract (n=194) | MSI | • Included in the data synthesis combining estimates from both bladder and upper urinary tract urothelial carcinomas  
o MSI was enriched in upper urinary tract (6.2% 12/194) relative to bladder cancer (0.9%, 4/454) with overall prevalence 2.5% (16/648) |
| Carlo (2019)35 | Kidney cancer | Renal cell carcinoma only | Metastatic | Apr. 2014 – Jan. 2017  
Total (N=115) | MSI | • Included in the data synthesis since this study focused on metastatic renal cell carcinoma only, which is the most common histologic sub-type of kidney cancer |
| Abida (2019)36 | Prostate cancer | | Overall | Jan. 2015 – Jan. 2018  
Total (N=1,551 from 1,346 patients) | MSI  
TMB (≥10 mut/Mb) | • Included in the data synthesis  
o Prevalence of MSI was reported separately for CRPC and non-CRPC: combined prevalence from both CRPC (4.5%, 16/356) and non-CRPC (2.4%, 16/677) cases were included in the data synthesis  
o Prevalence of TMB (≥10 mut/Mb) separately for CRPC and non-CRPC was not reported, and the overall prevalence was included in the data synthesis |
| Liu (2020)37 | Ovarian cancer | Advanced (mostly recurrent III/IV) | | Jan. 2013 – Apr. 2019  
Total (N=64) | MSI  
TMB (≥10 mut/Mb) | • Included in the data synthesis since the study included advanced ovarian cancer cases only |
| Stasenko (2020)38 | Endometrial cancer | Endometrioid carcinoma | IA | Jan. 2009 – Feb. 2017  
Total (N=211) | dMMR | • Included in the data synthesis since this study focused on stage IA endometrial carcinoma only, which is the most common histologic sub-type of endometrial cancer |

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; CRC = colorectal cancer; EC = endometrial cancer; IHC = immunohistochemistry; GE junction = gastroesophageal junction; CRPC = castration-resistant prostate cancer

a Number of samples available for the biomarker status.
4) Studies reporting the prevalence of dMMR/MSI/high TMB based on data from the Dana-Farber Cancer Institute

| Author (year) | Cancer(s)                  | Cancer sub-group(s) | Cancer stage(s)          | Data collection period and sample size* | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion                                                                 |
|---------------|----------------------------|---------------------|--------------------------|----------------------------------------|-------------------------|-------------------------------------------------------------------------------------------------|
| Albayrak (2020) | Pan-cancer                | 50+ solid tumours   | Overall                  | Aug. 2013 – July 2018                  | dMMR                    | Included in the data synthesis after excluding cancer types that do not satisfy the minimum sample size threshold <br>○ Haematologic tumours, lymphomas and benign lesions were also excluded <br>○ Prevalence predicted by an algorithm was included in the data synthesis given high concordance between the algorithm-based prevalence and the historical reports by pathologists (n=4,404) |
| Doyle (2019)   | Sarcoma                   | Soft tissue sarcoma | Overall                  | Period (N/S)                           | dMMR                    | Exclude from the data synthesis since this study cohort is a subset of the cohort reported in Albayrak et al. <br>○ Focused on reporting the different prevalence of dMMR between unclassified sarcomas (4/40, 10.0%) and classified sarcomas (3/264, 1.1%) |
| Christakis (2019) | Upper GI cancers  <br> Biliary tract cancers | Cancers in the <br> Small bowel <br> Stomach <br> Esophageal <br> Pancreas <br> Bile duct <br> Gallbladder <br> Ampulla | Overall                  | Period (N/S)                           | dMMR                    | Prevalence in esophageal (incl. gastroesophageal junction cancers) only was included in the data synthesis <br>○ Albayrak et al. reported the prevalence in esophagogastric cancers including both gastric cancers and esophageal cancers <br>○ Sample size of stomach cancer in this study does not satisfy the minimum sample size threshold (400+) <br>○ Prevalence in other cancer types was excluded from the data synthesis due to the smaller sample size than Albayrak et al. |
| Nassar (2019)  | Bladder/urothelial cancer | Urothelial carcinoma only | Overall <br> Stage-specific | 2013 – 2017 <br> Total (N=310) <br> Upper urinary tract (n=53) <br> Bladder (n=257) | TMB (≥10 mut/Mb) <br> TMB (≥20 mut/Mb) | Included in the data synthesis since Albayrak et al. did not report the prevalence of high TMB (≥10 mut/Mb or ≥20 mut/Mb) <br>○ A total of 162 T0 cases were excluded from the data synthesis |

\(^{a}\) Number of samples available for the biomarker status.

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified.
5) Studies reporting the prevalence of dMMR/MSI/high TMB based on data from Caris Life Science

| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|---------------|-----------|---------------------|-----------------|---------------------------------------|------------------------|----------------------------------|
| Nikanjam (2020)<sup>43</sup> | Pan-cancer | 40 tumour types | Overall | • Feb 2015 – Nov 2017  
• Total (N=28,034, excl. haematologic tumours, lymphomas ad benign tumours)  
  o MSI (n=28,034)  
  o TMB (n=27,847) | • MSI  
• TMB (≥17 mut/Mb) | • Prevalence of MSI was included in the data synthesis, excluding haematologic tumours, lymphomas ad benign tumours  
  o Pan-cancer prevalence  
  o Cancer-specific prevalence for those satisfying the minimum sample size threshold, except for the following cancer types  
    ▪ Biliary tract cancer: Spizzo et al.<sup>44</sup> reported the prevalence with the bigger sample size  
    ▪ Male genital tract malignancy, female genital tract malignancy: neither cancer-specific nor tumour group-specific given prostate cancer and ovarian cancer were reported, respectively  
    ▪ Uveal melanoma: this is a rare subtype of skin cancer  
• Prevalence of high TMB was included in the data synthesis in pan-cancer setting only due to the uncommon high TMB cut-off  
• No. of tumours with MSI and high-TMB were calculated based on the total no. of tumours and the reported prevalence in each cancer type (≥17 mut/Mb) |
| Spizzo (2020)<sup>44</sup> | Biliary tract cancer | Overall | June 2014 – Jan. 2019  
• Total (N=1,292) | • MSI  
• TMB (≥17 mut/Mb) | • Prevalence of MSI was included in the data synthesis due to the larger sample size than Nikanjam et al.<sup>43</sup>  
• Prevalence of high TMB was excluded from the data synthesis in pan-cancer setting only due to the uncommon high TMB cut-off |
| Tokunaga (2019)<sup>45</sup> | Appendiceal cancer | Adenocarcinoma | Overall | Apr. 2015 – Jan. 2018  
• Total (N=183) | • MSI  
• TMB (≥17 mut/Mb) | • Prevalence of MSI was included in the data synthesis  
  o Appendiceal cancer was not reported in Nikanjam et al.<sup>43</sup>  
• Prevalence of high TMB was excluded from the data synthesis due to the uncommon high TMB cut-off |
| Cimic (2020)<sup>46</sup> | Cervical cancer | • NECC  
• SCC | Overall | • Period N/S  
• Total (N=661)  
  o NECC (n=62)  
  o SCC (n=599) | • MSI  
• TMB (≥17 mut/Mb) | • Prevalence of MSI, combined both NECC (0/31, 0%) SCC (6/599, 1.0%), was included in the data synthesis  
  o Cervical cancer was not reported in Nikanjam et al.<sup>43</sup>  
• Prevalence of high TMB was excluded due to the uncommon high TMB cut-off |
| Author (year) | Cancer(s) | Cancer sub-group(s) | Cancer stage(s) | Data collection period and sample size$^a$ | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|---------------------|----------------|------------------------------------------|------------------------|----------------------------------|
| Jones (2020)$^{47}$ | Endometrial cancer | • Endometrioid carcinoma | • Overall | • Period N/S  
• Total (N=621) | • MSI  
• TMB ($\geq$10 mut/Mb) | • Prevalence of both MSI and TMB ($\geq$10 mut/Mb) were included in the data synthesis  
○ Endometrial cancer was not reported in Nikanjam et al.$^{43}$ |
| Seeber (2020)$^{48}$ | Pancreatic cancer | Pancreatic ductal adenocarcinoma | • Overall | • Apr. 2015 – Jan. 2018  
• Total (N=2,818) | • dMMR/MSI  
• TMB ($\geq$10 mut/Mb) | • Excluded from the data synthesis  
○ Prevalence of either dMMR or MSI was 1.3% in the tested tumours, but the number of tested tumours were not provided  
○ Nikanjam et al.$^{43}$ reported the similar prevalence of MSI (1.4%, 18/1261), and was included in the data synthesis |
| Stein (2019)$^{49}$ | Lung cancer | • NSCLC | • Advanced | • 2015-2017  
• Total (N=3,424) | • TMB ($\geq$10 mut/Mb) | • Included in the data synthesis  
○ Prevalence of high TMB was included in the data synthesis combining the prevalence from both HR-MT (28.3%, 230/812) and HR-WT (19.4%, 728/3750) |
| Heeke (2020)$^{50}$ | Breast cancer | • HR-MT  
• HR-WT | Overall | • Feb. 2015 - Jan. 2019  
• Total (N=4,562)  
○ HR-MT (n=812)  
○ HR-WT (n=3,750) | • dMMR/MSI  
• TMB ($\geq$10 mut/Mb) | • Prevalence of dMMR/MSI was excluded from the data synthesis  
○ Multiple test platforms were used to measure dMMR/MSI including fragment analysis, IHC and NGS and the combined prevalence of dMMR/MSI (0.6%, 26/4562) was reported.  
○ Nikanjam et al.$^{43}$ reported the similar prevalence of MSI measured by selected gene panel sequencing (0.7%, 17/2427) and was included in the data synthesis |

$^a$ Number of samples available for the biomarker status.

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; NECC = neuroendocrine cervical carcinoma; SCC = squamous cell carcinoma; NSCLC = non-small cell lung cancer; HR-MT = Homologous recombination DNA damage repair pathway mutated; HR-WT = Homologous recombination DNA damage repair pathway wild-type.
### Systematic reviews and/or meta-analyses reporting the prevalence of dMMR/MSI/high TMB

| Author (year) | Cancer(s) | Cancer stage(s) | Data collection period and sample size* | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|----------------|----------------------------------------|-------------------------|----------------------------------|
| Lorenzi (2020)51 | • Colorectal cancer  
• Endometrial cancer  
• Ovarian cancer  
• Gastric cancer  
• Esophageal cancer | • dMMR: Overall  
• MSI  
  o Overall  
  o I/II, III/IV (subgroup) | • ~ Oct. 2017 | • dMMR  
• MSI | • Pooled overall and stage-specific prevalence of dMMR and MSI across all tumours reported in this structured/targeted review were not included in the data synthesis  
• Overall cancer-specific prevalence of dMMR (ovarian cancer, gastric cancer) and prevalence of MSI (ovarian cancer, gastric cancer, and esophageal cancer) only were included in the data synthesis due to different stage grouping from this scoping review  
• Prevalence of dMMR and MSI in colorectal cancer and endometrial cancer were excluded from the data synthesis  
  o Colorectal cancer (dMMR: 13.2%, 1513/11434; MSI: 11.5%, 937/8156): prevalence estimates were obtained through targeted review, and a systematic review by Jin et al.52 was included in the data synthesis  
  o Endometrial cancer (dMMR: 24.8%, 1302/5248; MSI: 26.0%, 1773/6813): Ryan et al.53 was the most up-to-date systematic review of endometrial cancer and the research questions aligns better with this scoping review, and this study was included in the data synthesis |}

| Luchini (2020)21 | Pancreatic ductal adenocarcinoma | Overall | • ~30/11/2019  
• Total (N=8,323 cases from 34 studies) | • dMMR/MSI  
  o MSI by NGS  
  o dMMR/MSI by IHC/PCR | • Prevalence of MSI measured by selected gene panel sequencing alone was included in the data synthesis  
  o Statistically significant difference in the prevalence by assays used: gene panel sequencing (1.1%, 68/6030) vs IHC/PCR (6.5%, 150/2293)  
  o Most included studies used PCR for MSI analysis was not with recommended panel of markers (nor NCI neither MSI PCR).  
  o Included studies often reported the combined prevalence of dMMR/MSI by IHC/PCR  
  o Of the included studies, three studies were published in 2019 (Latham et al.28, Singhi et al.,20 and Kato et al.54), of which the study period overlaps with this scoping review. All three studies were also identified in this scoping review, and the prevalence estimates from these original research studies were not included in the data synthesis |
| Author (year) | Cancer(s) | Cancer stage(s) | Data collection period and sample sizea | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|----------------|----------------------------------------|-------------------------|----------------------------------|
| Ryan (2019)  | Endometrial cancer | Overall | ~ July 2018  
Total (N=12,633 cases from 53 studies) | dMMR  
MSI | Included in the data synthesis  
○ Prevalence of dMMR  
○ Prevalence of MSI |
| Kahn (2019)  | Endometrial cancer | dMMR  
I, II, III, IV (sub-group)  
MSI | Jan. 1990 - Jan. 2018  
Total (N=6,649 cases from 29 studies)  
○ dMMR (n=6,649)  
○ MSI (n=3,140) | dMMR  
MSI | Prevalence of dMMR in early-stage endometrial cancer only was included in the data synthesis  
○ Likely to be a subset of studies included in a systematic review by Ryan et al.53  
○ Prevalence in advanced-stage endometrial cancer does not satisfy the minimum sample size threshold (n=24) |
| Jin (2020)   | Colorectal cancer | Overall | 2007 – July 2018  
Total (N=17,621 from 44 studies) | dMMR  
MSI | Included in the data synthesis due to separate reporting of the prevalence of dMMR and MSI, although the sample size was smaller than John et al.56 |
| John (2020)  | Colorectal cancer | Overall | 2005 – 2017  
Total (N=47,545 from 73 studies) | dMMR/MSI | Included in the data synthesis for the combined prevalence of dMMR and MSI  
○ High concordance between dMMR and MSI in colorectal cancer  
○ Research question is aligned with this scoping review: Systematic review of studies performing universal screening for LS |
| Wang (2019)  | Colorectal cancer | III  
IV | ~ July 2018  
Total (N=21,175 from 36 studies)  
○ Stage III (n=18,277)  
○ Stage IV (n=2,898) | dMMR/MSI | Included in the data synthesis for the combined prevalence of dMMR and MSI  
○ Stage-specific prevalence was reported  
○ High concordance between dMMR and MSI in colorectal cancer |
| Deng (2020)  | Colorectal cancer | II/III combined | ~ May 2019  
Total (N=28,331 from 51 studies) | dMMR | Included in the data synthesis for early-stage colorectal cancer  
○ Of the included studies, only one study was published in 2019 (Fountzilas et al.59), of which the study period overlaps with this scoping review. This study was also identified in this scoping review and the prevalence estimate from this original study was excluded from the data synthesis |
| O’Connell (2020) | Rectal cancer | II/III combined | ~ Aug. 2019  
Total (N=5,877 from 9 studies) | dMMR/MSI | Included in the data synthesis for the combined prevalence of dMMR and MSI  
○ Stage-specific prevalence was reported  
○ High concordance between dMMR and MSI in colorectal cancer |
| Author (year) | Cancer(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|----------------|---------------------------------------|------------------------|----------------------------------|
| Willis (2019)<sup>62</sup> | Lung cancer | Advanced | Jan. 2012 ~ Apr. 2018 | TMB (≥10 mut/Mb) | Included in the data synthesis from 3 studies reported the prevalence of high TMB using a common high TMB cut-off (≥10 mut/Mb) <br> o Two NSCLC studies: Checkmate 026, Checkmate 227 <br> o One SCLC study: Checkmate 032 |
| Zhu (2019)<sup>63</sup> | Lung cancer | Advanced | ~ Oct. 2018 | TMB (≥10 mut/Mb) | Excluded from the data synthesis <br> o Of the included studies, only two studies used the common high TMB cut-off TMB (≥10 mut/Mb or 200+ mutations from WES), and both of them were NSCLC studies that were also included in Willis et al.<sup>62</sup> |

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; IHC = immunohistochemistry; PCR = polymerase chain reaction; LS = Lynch syndrome; NSCLC = non-small cell lung cancer; SCLC = small cell lung cancer.

<sup>a</sup> Number of samples available for the biomarker status.
| Author (year)      | Cancer(s) | Cancer stage(s) | Data collection period and sample size | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------------|-----------|----------------|---------------------------------------|--------------------------|---------------------------------|
| Salem (2020)⁶⁴     | Colon cancer | III            | • 1998 - 2009  
• Total (N=6,501) | dMMR/MSI | • Included in the data synthesis for the combined prevalence of dMMR and MSI given high concordance between dMMR and MSI in colorectal cancer  
○ Pooled analysis of patients from six adjuvant CT trials (MOSAIC, C07, C08, PETACC8, N0147, AVANT) treated with fluorouracil, leucovorin, and oxaliplatin and included in the ACCENT database  
○ Larger sample size than Taieb et al.⁶⁵ |
| Taieb (2019)⁶⁵     | Colon cancer | III            | • 1998 - 2009  
• Total (N=2,630) | dMMR/MSI | • Excluded from the data synthesis due to the smaller sample size than Salem et al.⁶⁴  
○ Pooled analysis of patients from six adjuvant CT trials (MOSAIC, C07, C08, PETACC8, N0147, AVANT) focused on those with disease recurrence following adjuvant treatment |
| Sinicrope (2021)⁶⁶  | Colon cancer | III            | • 2004 - 2009  
• Total (N=5,337) | dMMR/MSI | • Excluded from the data synthesis  
○ A subset of six adjuvant CT trials (MOSAIC, C07, C08, PETACC8, N0147, AVANT) included participants from PETACC8 and N0147 only |
| Pietrantonio (2019)⁶⁷  | Gastric cancer | II/III  | • Period (N/S?)  
• (N= 1,556 from 4 RCTs) | MSI | • Included in the data synthesis  
○ Meta-analysis of individual patient data from four RCTS (MAGIC, ITACA-S, ARTIST, CLASSIC) compared surgery with surgery + CT for resectable gastric cancer in four countries |
| Choi (2019)⁶⁸       | Gastric cancer | II/III  | • Period (N/S?)  
• Total (N= 592) | MSI | • Excluded from the data synthesis  
○ A subset of the study cohort reported in Pietrantonio et al.,⁶⁷ including participants from CLASSIC only |
| Barroso-Sousa (2020)⁶⁹  | Breast cancer | Overall  | • Period (N/S?)  
• Total (N=3,951)  
○ Primary cancer (n=2,455)  
○ Metastatic cancer (n=1,496) | TMB (≥10 mut/Mb) | • Included in the data synthesis  
○ Genomic and clinical datasets from three WES studies and three targeted panel studies, including GENIE-DFCI-ONCOPANEL-3, GENIE-MSK IMPACT410, and GENIE-MSK IMPACT468.  
○ Original research study from the MSK breast cancer cohort was not identified in this scoping review.  
○ One original research study from the DFCI breast cancer cohort was identified in this scoping review and the prevalence of dMMR reported in Albayrak et al.³⁹ was included in the data synthesis |
| Author (year) | Cancer(s) | Cancer stage(s) | Data collection period and sample size<sup>a</sup> | Pan-tumour biomarker(s) | Rationale for inclusion/exclusion |
|--------------|-----------|----------------|--------------------------------|-------------------------|---------------------------------|
|              |           |                |                                |                         | ○ Overall prevalence using all the samples (50%, 196/3951) and the prevalence in advanced breast cancer using samples from metastatic cancers only (8.4%, 125/1496) |

dMMR = mismatch repair deficiency; MSI = microsatellite instability; TMB = tumour mutational burden; N/S = not specified; CT = chemotherapy; RCT = randomised controlled trial; WES = whole exome sequencing; MSK = Memorial Sloan Kettering Cancer Centre; DFCI = Dana-Farber Cancer Institute.

<sup>a</sup> Number of samples available for the biomarker status.
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