Comparing nodal versus bony metastatic spread using tumour phylogenies

Stefano Mangiola1,2, Matthew K. H. Hong1, Marek Cmero1,2, Natalie Kurganovs1, Andrew Ryan3, Anthony J. Costello1,4, Niall M. Corcoran1,4, Geoff Macintyre2,5 & Christopher M. Hovens1,4

The role of lymph node metastases in distant prostate cancer dissemination and lethality is ill defined. Patients with metastases restricted to lymph nodes have a better prognosis than those with distant metastatic spread, suggesting the possibility of distinct aetiologies. To explore this, we traced patterns of cancer dissemination using tumour phylogenies inferred from genome-wide copy-number profiling of 48 samples across 3 patients with lymph node metastatic disease and 3 patients with osseous metastatic disease. Our results show that metastatic cells in regional lymph nodes originate from evolutionary advanced extraprostatic tumour cells rather than less advanced central tumour cell populations. In contrast, osseous metastases do not exhibit such a constrained developmental lineage, arising from either intra or extraprostatic tumour cell populations, at early and late stages in the evolution of the primary. Collectively, this comparison suggests that lymph node metastases may not be an intermediate developmental step for distant osseous metastases, but rather represent a distinct metastatic lineage.

It has long been recognized that the spread and seeding of metastatic tumour cells from the primary site of origin does not follow a random pattern, rather, there are characteristic anatomical sites favouring the formation of metastatic lesions1. For epithelial tumours such as prostate adenocarcinoma, the bony skeleton as well as regional lymph nodes are a common site of metastatic spread. Depending on the tumour stage and grade, as well as the anatomical extent of dissection, lymph node metastases can be identified in up to 8% of patients undergoing radical prostatectomy in contemporary series2–4. However the clinical behaviour of lymph node metastases differs from that of bony and other visceral disease. Patients presenting with lymph node only metastases have significantly better progression-free and overall survival compared to patients with mixed lymph node and bony disease, bony disease alone, or spread to other visceral sites, and consistently demonstrate an improved therapeutic response to hormonal manipulation and chemotherapy5. In addition, in a randomised study evaluating the impact of early androgen deprivation in patients with N1 disease, 14% of patients were disease free at a median of 11.9 years postoperatively with no further treatment, suggesting at least the possibility of long term disease control if not cure with surgical excision alone6.

For over a century, lymph nodes have been considered to have an essential role in the metastasis of epithelial tumours2, acting as a staging ground for wider visceral dissemination8. Recent clinical studies however suggest that metastatic spread is a more complex process, and that tumour colonisation of regional lymph nodes may not directly contribute to lethality, but instead represent an evolutionary ‘cul-de-sac’ that may be a pathological marker of inherently aggressive disease but does not play a direct role in systemic colonisation9. Importantly it remains to be clarified whether visceral metastases arise from previously established metastatic lymph nodes or if lymph nodes are irrelevant to the evolution of distant metastases. The understanding of the role of lymph node metastases in prostate cancer evolution will provide key insights into the mechanisms leading to the lethal disease.
Recent studies in prostate cancer\cite{10,11} used tumour phylogenies to reveal the complexity of distant tumour spread, describing patterns of cross metastatic seeding, multiple metastatic waves of spread from the primary, and a possible metastatic origin of local recurrences.

Several evolutionary models that attempt to describe the dynamic processes driving tumour cell diversification and spread to distant organs have been proposed in the pre-next-generation sequencing era, including clonal evolution\cite{12} the mutator phenotype\cite{13,14} and stochastic progression\cite{14}. Using next-generation sequencing data, algorithms that reconstruct phylogenies of metastatic cancers based on nucleotide variants have been recently developed\cite{15}. Tumour phylogenetic methods that permit the tracing of tumour cell lineages in human cancers have been used to identify the roots of metastasis in humans\cite{16} and, along with mapping in a comprehensive manner the sub-clones within the primary tumour, has been stressed to be key for the reconstruction of the metastatic spread\cite{17}. However, these studies did not characterise localised lymph node metastases.

To address this, we collected a total of 19 samples from multiple prostate tumour foci from 3 patients, together with matched metastatic lymph nodes. We performed genome-wide single nucleotide polymorphism (SNP) array profiling of each sample and characterised all copy-number changes. We then combined these data with 26 samples from 3 men with distant metastatic disease\cite{18}. In order to identify the patterns of copy number aberrations (CNAs) that emerged at the transition across histo-pathological categories, a combined evolutionary reconstruction and differential analysis of gene copy number was performed. Using these copy-number profiles we were able to derive tumour phylogenetic trees for each patient using MEDDIC\cite{19}, combined with a bootstrapping procedure to refine and improve robustness of the trees. Each tree was then validated using targeted Multiplex Ligation-dependent Probe Amplification (MLPA) of 58 genomic regions spanning 39 genes. Analysis of the tumour phylogenies revealed distinct patterns of metastatic spread between the two patient groups, as well as potential drivers of transition across histo-pathological categories. We show lymph node metastases originated exclusively from evolutionarily advanced, extraprostatic regions of primary tumour, whereas distant metastases originated either from central (2 cases) or extraprostatic primary tumour regions (1 case).

**Results**

**A comparison of broad copy-number changes between tumour locations identifies shared and unique events.** We used the Illumina BeadChip to determine the genome-wide copy-number profiles of 48 samples collected from 6 patients with metastatic prostate cancer (see Methods section). 37 were inferred as approximately diploid, 10 were found with to have an average ploidy close to 3, and 1 sample had a tetraploid genome (Fig. 1A, Table S1). Primary and metastatic disease sites were found to have examples of both diploid and non-diploid genomes.

We grouped the samples by histo-pathological category (central, extraprostatic, lymph node, and bony metastasis), adjusted for ploidy, and computed a summary copy-number profile (Fig. 1B). Visual inspection of the profiles indicated shared changes across the groups: recurrent 8p deletions (8p22 to 8p11.23) and 8q amplifications (8q11.21 to 8q24.23), were observed similar to previous studies\cite{20}, along with deletion of the long arm of chromosome 16 (16q22.1 to 16q24.3)\cite{21}. Given these changes are shared across groups, they are likely to have occurred multiple times through evolution of each tumour.

The only region to show marked divergent frequencies between groups was 4q (Fig. 1B), with primary tumour samples showing no change, lymph node metastases showing amplification and bony metastases showing deletion. This region harbours TET2, previously shown to be deleted in metastatic disease\cite{22} and its expression linked with disease progression\cite{23}.

**Tumour phylogenies reveal the origin of lymph node metastases as exclusively extraprostatic.** In order to identify clones and infer their evolutionary hierarchy in each patient we used MEDDIC\cite{19} in combination with a bootstrapping strategy to improve tree robustness (see Methods section). The resulting phylogenies (Fig. 2) show that lymph node metastases have a recurrent evolutionary pattern originating exclusively from regions of extraprostatic extension (EPE), never directly from clones present within the central prostatic capsule. In contrast, the distant osseous metastatic clones originated from either central or extraprostatic locations. Central origin can be seen in patients 001 and 498 where no extraprostatic, localized tumour was identified upon pathological review, either in the SV or EPE. In these two cases all of the distant metastases (i.e., C, B and G for patient 001; and MC, R, MFT and MA for patient 498) originated from an ancestor (i.e., ii and iii respectively) derived from a tumour locus localized in a deep, central position in the prostate suggesting an early tumour population located extraprostatically in the seminal vesicle (i.e., K)\cite{24}.

In order to validate these observations we used an orthogonal copy-number profiling approach, Multiplex Ligation-dependent Probe Amplification (MLPA) technology. We processed samples C, D, G, H, J and L from patient 167; C, D, E, G and L from patient 179; and C, E, F, G, H and L from patient 421. For the selected samples we tested a total of 39 genes, utilizing a total of 63 different probes. Overall, we validated the direction of variation in copy number (i.e., deletion, amplification or diploid) for over 71% of the probes across all selected samples. For each branch of the evolutionary hierarchies, we validated an average of 68% of the probes for the genes that define that branching (Table 1). Although MLPA technology was not designed for taking into account samples with highly variable tumour purity, we could obtain a validation p-value <10^-4 for each branch along the evolutionary schema (Table 1), using a binomial test (binom.test function in R; using 0.33 for the p parameter, indicating three possible categorical outcomes for copy number changes: decrease, increase, no-change). The evolutionary hierarchies for patients 001, 177, 299 and 498 were consistent with previous analyses based on whole genome sequencing data\cite{25}. For patient 001 (Fig. 3), both our analyses and published data indicate that the distant metastases had a common central tumour ancestor, namely, tumour sample C. Similarly for patient 498, the distant metastases consistently clustered together having a single common ancestor derived from a central primary tumour. For
patient 299, the relationship between the various metastatic and primary tumour foci for this patient was derived from the WGS data, due to biases in the SNP array data as described.

**Tumour phylogenies facilitate identification of copy-number changes associated with metastatic progression.** We used the inferred clonal phylogenies to associate specific copy-number events with the transition from diploid-to-central tumour, central-to-EPE, EPE-to-lymph node metastasis, and central/EPE-to-bony metastasis (see Methods, Fig. 3, and Table 2).

A number of interesting observations can be made from this analysis including refinements of the broad copy-number changes observed earlier. The analysis shows that 8p deletions are observed at low frequencies across all four transitions, albeit at slightly higher frequencies in the transition to bony metastases, suggesting a potential necessary condition for spread to bone. Broad 8q gains appear at high frequency during the emergence of the central tumour, however, in the transition to EPE, there is a focal deletion (within the base pair interval 77796946–116276618). This likely represents a loss-of-heterozygosity that confers ability to spread and invade (Fig. 3). Further amplification of 8q is observed in both transitions to lymph and bone metastases, likely as a result of the necessity to increase c-Myc gene copy-number, an important regulator of proliferation.

16q deletions were detected as significantly altered, at low frequency in central tumour development and metastasis to bone, and at high frequency in the development of lymph node metastases. This region includes FBXO31, a key oncosuppressor in prostate cancer that commonly displays loss of heterozygosity, including in breast, ovarian and hepatocellular cancers. This alteration may be necessary for the ability to metastasise from the EPE to lymph nodes. Two other loci in chromosome 16 also show differential behaviour between categories (Table 2): 16p13.3 from chromosome 16, which contains CBP, the transcriptional co-activator of CREB (cAMP response element binding protein); this gene regulates cell differentiation, proliferation and survival and is involved in prostate tumour initiation. Additionally, the locus 16q12 from chromosome 16 had a decrease in copy number during the transition to lymph node. This region includes two key genes: CYLD and BRD7. The gene CYLD is a deubiquitination enzyme which regulates cell survival or cell proliferation, and is commonly lost in different types of human cancer; the gene BRD7 negatively regulates cell proliferation and growth and has a key role in prostate cancer development.

Further alterations can be found in 11p15.5 from chromosome 11, which is decreased in both the transition to extraprostatic tumour and to lymph node metastases. This region includes MUC2, a known tumour suppressor; and MUC5, which have been previously described as under-expressed in prostate cancer.
The gene TET2 located in the locus 4q24 increased in copy number with higher frequency in the transition to central tumour and lower frequencies in the transition to lymph node and bony metastases. Finally, three other loci showed altered copy numbers: 4q24, 5p15.2 and 19q13.4, containing potential drivers in NPNT, SEMA5A and BRSK1 respectively.

Global copy-number profile comparison identifies lymph and bony metastases as distinct disease states. Finally, we sought a measure to determine the genome-wide relatedness between lymph, bone, EPE and central tumours that utilised the tumour phylogenies. CNA profiles belonging to the same histo-pathological category were averaged, obtaining four centroid profiles for each category. We then calculated the Euclidean distance between each applying a bootstrap leave-one-out method that produced seven distinct distance matrices, which showed consistent results to the predicted solution in Fig. 4.

Table 1. Validation using MLPA. Significance calculated with binomial test with \( p = 0.33 \). P-values were corrected for multiple test with the Benjamini, Hochberg, and Yekutieli method\(^\text{44}\).

| Patient | Node | descendants | Probes validated (%) | P-value | Adjusted p-value |
|---------|------|-------------|----------------------|---------|-----------------|
| 167     | i    | ii; iii     | 76.3                 | 2.86e-11 | 1.7178e-10      |
| 167     | iii  | iv; D       | 73.7                 | 2.55e-10 | 6.1224e-10      |
| 167     | iv   | v; L/H      | 70.0                 | 7.34e-09 | 1.4686e-08      |
| 179     | i    | ii; vi      | 73.7                 | 2.55e-10 | 6.1224e-10      |
| 179     | ii   | iii; v      | 73.7                 | 2.55e-10 | 6.1224e-10      |
| 179     | iii  | iv; G       | 87.5                 | <2.2e-16 | 2.64e-15        |
| 421     | i    | ii; B       | 60.7                 | 3.10e-05 | 3.7248e-05      |
| 421     | ii   | iii; D      | 60.7                 | 3.10e-05 | 3.7248e-05      |
| 421     | iii  | iv; C       | 60.7                 | 3.10e-05 | 3.7248e-05      |
| 421     | iv   | v; H        | 61.3                 | 7.81e-06 | 1.3404e-05      |
| 421     | v    | iv; F       | 57.1                 | 2.15e-04 | 0.0002159       |
| 421     | vi   | vii; L      | 59.0                 | 6.76e-05 | 7.378909e-05    |

Figure 2. Cancer evolutionary hierarchies for patients 167, 179, 421 (including lymph node metastasis) and 001, 299, 498 (including distant metastasis). The roman numbers indicate the predicted ancestral cell populations. The decimal numbers at branches represent the confidence values of the clade prediction (when smaller then 1).
The analysis of the Euclidean distance across copy-number profiles (Table S2) surprisingly indicated that the profiles of bone metastases were generally more related with the primary prostate tumour populations than with the lymph node metastases. The large separation between lymph and bone metastases using this measure suggests they are distinct at a genomic level.

Discussion
In this study we explored the genomic relationships between distinct metastatic populations in a small number of advanced prostate cancer patients. Inspection of the evolutionary hierarchies that we derived for patients with lymph node metastases, 167, 179 and 421, indicated that these metastases most likely originated from tumour cell populations that had already extended beyond the prostate capsule and which therefore exhibited an invasive phenotype, rather than from more centrally located and less invasive tumour cell populations, such as in the case for distant bone metastases. This suggests that lymph node tumours may not act as a conduit point for further tumour cell dissemination, at least not to more distant osseous or visceral sites. These observations are also consistent with the model of metastatic progression that emerges from a similar study of advanced prostate cancer patients. In this study, the evolutionary hierarchies that include lymph node metastases (i.e., patients A17, A22, A24, A31 and A32) consistently show that lymph node metastasis is not a common ancestor for distant bone metastases, but rather represents an independent evolutionary route emerging from central tumours. Furthermore, a recent study for brain metastases identified regional lymph node metastases as highly divergent from the metastases located in the brain; supporting the hypothesis of diverse evolutionary histories of local or distant metastases.

These results, though not by any means definitive due to the limited sample size, do seem to accord with the observed differences in clinical behaviour of different anatomical sites of metastasis in advanced prostate cancer patients. Prior clinical studies documenting metastatic status and oncological outcomes in prostate cancer patients have lent weight to the hypothesis that lymph node metastasis is a distinctive biological entity compared to metastases presenting at bony or visceral sites. For instance it has been demonstrated that there is a clear differential in the survival status of patients with node only disease even after primary treatment, compared to those with more distant metastases. More recent studies are now also corroborating that the survival rates of patients presenting with metastatic disease is significantly linked to both the site and number of metastatic deposits. Individuals from a large contemporary cohort of prostate cancer patients presenting with either exclusive visceral metastases or additionally with bone involvement, had poorer survival than those with bone only metastases.

Figure 3. Loess regression of the increase/decrease CNA profiles, obtained from the difference in copy numbers across transition between predicted ancestral representative of the four histo-pathological categories (i.e., from diploid-to-central tumour, central-to-EPE, EPE-to-lymph node metastasis, and central/EPE-to-bony metastasis) for each patient (one per patient, per category). The value of 0 of the y axes represents no variation compared to the parent histo-pathological category. The light-blue shades represent the genomic regions having a differential amount of copies across the four transitions (p-value < 0.1, after method Benjamini, Hochberg, and Yekutieli multiple test correction).

The analysis of the Euclidean distance across copy-number profiles (Table S2) surprisingly indicated that the profiles of bone metastases were generally more related with the primary prostate tumour populations than with the lymph node metastases. The large separation between lymph and bone metastases using this measure suggests they are distinct at a genomic level.
Table 2. Genes within the top 10 regions for differential gain/loss copy number aberration.

| Chromosome | Locus | Gene     | Copy number increase/decrease (mean) | P-value | Adjusted p-value |
|------------|-------|----------|--------------------------------------|---------|-----------------|
| 16         | 16q24.2| FBXO31   | Central: −0.33, Extraprostatic: 0.25, | 0.01    | 0.0006          |
|            |       |          | Lymph node: −1.67, Bone: 0.01        |         | 0.0312          |
| 11         | 11p15.5| MUC2     | 0                                    | 0.69    | 0.0033          |
|            |       |          | −0.5                                |         | 0.08013         |
| 11         | 11p15.5| MUC6     | 0                                    | 0.69    | 0.0033          |
|            |       |          | −0.5                                |         | 0.08013         |
| 11         | 11p15.5| KRTAP5-5 | 0                                    | 0.69    | 0.0033          |
|            |       |          | −0.5                                |         | 0.08013         |
| 1          | 1p36.33| CFA7P74  | 0                                    | 0.41    | 0.0157          |
|            |       |          | −0.67                               |         | 0.11452         |
| 16         | 16p13.3| CREBBP   | 0                                    | 0.5     | 0.0175          |
|            |       |          | 0.25                                |         | 0.11452         |
| 8          | 8q24.3 | GML      | 0.8                                  | 0.54    | 0.0207          |
|            |       |          | −0.67                               |         | 0.11452         |
| 8          | 8q24.3 | MAF      | 0.8                                  | 0.54    | 0.0207          |
|            |       |          | −0.67                               |         | 0.11452         |
| 4          | 4q24   | TET2     | 0.67                                 | 0.33    | 0.0385          |
|            |       |          | −0.5                                |         | 0.18741         |
| 4          | 4q24   | NPNT     | 0.67                                 | 0.33    | 0.0385          |
|            |       |          | −0.5                                |         | 0.18741         |
| 19         | 19p13.4| BRSK1    | 0.17                                 | 0.51    | 0.0497          |
|            |       |          | 0.25                                |         | 0.18741         |
| 8          | 8q24   | RAD21    | 1.4                                  | 0.5     | 0.0604          |
|            |       |          | −0.67                               |         | 0.21143         |
| 5          | 5p15.2 | SEMA5A   | 0                                    | 1.01    | 0.0701          |
|            |       |          | 0.5                                 |         | 0.21206         |
| 8          | 8q23.1 | ANGPT1   | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22   | RUNX1T1  | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22.2 | COX6C    | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21   | NBN      | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22.3 | RIMS2    | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22.2 | CPQ      | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21   | WWP1     | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21   | HEY1     | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22.2-q23| PABPC1 | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22   | UBR5     | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21.13| PAG1     | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q23   | DCSTAMP  | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q23   | PHD1L1   | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q23.3 | CSMD3    | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21.3 | DCAF4L2  | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q21.3 | SLC7A13  | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q22.3 | SNX31    | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |
| 8          | 8q23.1 | RSPO2    | 1.6                                  | 0.5     | 0.0741          |
|            |       |          | −0.67                               |         | 0.21206         |

Figure 4. Multi-dimensional scaling plot of the centroids of the increase/decrease copy number profiles representative of the transitions between the four histo-pathological categories (i.e., from diploid-to-central tumour, central-to-EPE, EPE-to-lymph node metastasis, and central/EPE-to-bony metastasis). Representative copy number profiles for distant metastases were taken from patients 001, 177, 299, 498; representative copy number profiles for the other histo-pathological categories were taken from patients 167, 179 and 421.
The number of metastatic sites involved also impacted on survival with patients having multiple metastatic sites having a significantly shorter survival than patients with oligometastatic disease. Interestingly, patients with a single visceral metastasis experienced a higher mortality rate than patients with bony metastatic disease. Patients however who exhibited only lymph node involvement had the best mortality outcomes of any of the metastatic patients and a significantly better overall and cancer specific survival than patients with visceral metastases. It is noteworthy that all the patients from this study had stage IV disease at presentation and furthermore the node only disease patients were only diagnosed using the relatively insensitive imaging modalities of CT and MRI and hence it is likely that their true nodal disease burden was in fact underestimated. It is tempting to speculate that the clear differences in oncological outcomes of these patients based on the location of the metastatic deposits reflects underlying biological differences in the nature of the tumour cell clones that give rise to the distinctive patterns of metastatic disease.

There has been very little genetic investigation of the potential underlying differences in metastatic lesions based on site of presentation in prostate cancer; however work in the field of breast cancer has shed some light on this issue. Data from preclinical modelling of breast cancer metastasis indicates that distinct populations of tumour cells in the primary organ drive spread to lymph nodes and that these cells possess distinctive epigenetic profiles that predispose them to disseminate and proliferate in the lymph nodes rather than bony sites. In breast cancer, patients with positive lymph nodes have a clearly increased risk of disease recurrence following lymphadenectomy surgery. Additionally, lymph node status is the single most important prognostic feature in breast cancer and in this regard the disease differs from prostate cancer. Recently, it has been demonstrated that differential methylation of the PAX6 gene was found to be linked to lymph node metastasis in breast cancer, with positive nodes having significantly lower levels of PAX6 methylation than paired and matching primary tumour tissues from the same cohort of 25 patients. These results further strengthen the hypothesis that specific tumour cell clones with distinctive epigenetic changes might disseminate and colonize specifically the lymph node niche in cancer patients.

A number of key questions are yet to be resolved concerning the mechanisms of distinctive tumour cell dissemination. For example, do cells with these distinctive genetic changes arise stochastically in unselected populations localized within the primary tumour or do metastasis specific changes occur first in the prostate tumour cells and then once the cells have lodged in specific anatomical locations are they then further selected for changes that predispose them to proliferate and colonize distinctive metastatic niches? Certainly our results suggest that the tumour cell populations that seed to either lymphatic or bony sites arise from distinct tumour locations in the prostate and may therefore be indicative of the biological characteristics of tumour cell population of origin as a whole.

Resolving the issue of whether positive lymph nodes can commonly act as a conduit for more distant bony or visceral tumour dissemination is not just of academic interest but has clear clinical implications. This is particularly so in the setting of the burgeoning uptake in the new and highly sensitive imaging modality of PSMA PET. With the increasing use of this screening technology an increasing number of node only as well as node plus osseous and/or visceral sites metastatic patients are likely to be identified.

This study represents a foundation for a further investigation of the behavioural patterns of prostate cancer metastases at distinct sites. Such improved knowledge will allow greater metastatic control as well as the employment of therapies that target selectively high risk lesions. This will result in decreased systemic treatment toxicity compared to current systemic therapies that assume all metastatic lesions are equivalent.

Methods

All experimental protocols and all methods described here were approved and carried out in accordance with the guidelines of the Epworth Hospital Human Ethics committee.

Patient selection. Following informed consent from all patients and institutional ethics approval from the Epworth Hospital Human Ethics committee, suitable prostate cancer specimens were identified from three patients with N1 disease following radical prostatectomy and lymph node dissection (i.e., 167, 179 and 421) from our institutional database. As a comparison arm, we included in the study a further four patients with biopsy confirmed bony metastatic disease following radical prostatectomy. All patients prospectively consented to the use of their tissues for genomics studies in accordance with the core bioethical elements for genomics projects. Clinico-pathological details of these three patients are presented in Table S1. All three patients presented with large, advanced stage, primary tumours. No patient had more than one lymph node histologically confirmed to contain malignancy. The prostate specimens were fixed in formalin after collection. Glands were sliced at 4–5 mm intervals perpendicular to the urethra, with the apex and base slices additionally sectioned in the parasagittal plane at 4 mm intervals. A 5 mm slice was then subsequently made from each section, stained with haematoxylin and eoxin (H&E) and all tumour borders marked by pen. These slides were then scanned and a tumour map constructed depicting the topography and extent of prostate cancer in each gland as well as any features of prognostic relevance, such as seminal vesicle involvement. All pathological examinations of specimens were undertaken prospectively and shared within a group of four urological pathologists with regular internal audit practices for consistency.

Tissue processing. We identified regions of interest based on topographical depiction of tumour location, extent and foci in relation to the rest of the prostate gland; and targeted areas of tumour that defined high-risk disease, including seminal vesicle invasion and extraprostatic extension, as well as adjacent central portions of tumour and other foci of lower grade cancer. We also included positive surgical margins as these areas could represent the advancing front of the tumour. For lymph nodes, we targeted tumour at the capsule of the lymph node at the marginal sinus, considering that this region is prone to early deposits of metastatic prostate cancer.

---

SCIENTIFIC REPORTS | 6:33918 | DOI: 10.1038/srep33918
We then cut 5 μm reference slide from archival blocks of formalin-fixed paraffin-embedded (FFPE) containing regions of interest. The slide was H&E stained and boundaries of tumour foci were marked with felt-tip pen on the cover slips. A further six slides of 10 μm thickness were cut from each block for macrorossection. The unstained slides were deparaffinised in xylene, washed with a decreasing ethanol series and nuclease-free water, and H&E stained with a rapid protocol. The slides remained immersed in ethanol until ready for dissection.

We used three to six slides per region of interest depending on the likely DNA yield; and macrorossected each region of interest using 27.5 G hypodermic needles and number 15 scalpel blades at 8–35× magnification (Leica EZ4 HD stereomicroscope).

**DNA Extraction and SNP Chip.** Genomic DNA was extracted using the QIAamp DNA FFPE Tissue Kit (Qiagen, Maryland, Ca) according to manufacturer’s instructions. Briefly, we placed the macrorossected tissue in 180 μl ATL buffer with 20 μl protease K, and performed RNase digestion and genomic DNA purification by column chromatography with elution in 100 μl nuclease-free water. We restored 200 ng of genomic DNA per sample using the Infinium HD FFPE Restore Protocol (Illumina) and Zymo Research DNA Clean & Concentrator (Zymo) as required. For each sample, all 8 μl of restored DNA was used as input for the Infinium HD FFPE Assay (Illumina). We linearly amplified the DNA across the whole genome and fragmented it enzymatically. The resulting product was hybridised to the Illumina HumanOmniExpress-FFPE-12 v1.0 BeadChip and incubated at 48°C for 16–24 hrs. Imaging was performed using the Illumina iScan system and intensity values derived for each bead type. LogR Ratios and B allele frequencies (BAF) were calculated from the intensity data using GenomeStudio v2010.3 with Genotyping module 1.8.4 software and the HumanOmniExpress-12v1 G FFPE manifest cluster file. Overall, we achieved high SNP call rates with a median of 98.35% (range 92.47% to 99.72%). The median standard deviation of the LogR Ratio values was 55%. The goodness of fit of the predicted aberrant fractions was always above 0.76 with a mean of 0.91.

**Phylogenetic analyses.** For the prediction of gene copy number (CN) profiles, tumour cellularity and ploidy of the patients 167, 179, 421 (for which sample collection, processing and molecular analysis is described above), and for patients 001, 177, 299, 298 (for which sample collection, processing and molecular analysis has been described previously) we employed ASCAT with default settings. For each of those patients, with the exception of patient 177 (for which no central tumour was available), we inferred the tumour evolutionary hierarchy from the copy number profiles (Fig. S1). First, we fragmented the genomic breakpoints (defined as the start and end of the genomic regions affected by CNA) of all samples to obtain comparable CNAs profiles, and then selected CNAs originated from large genomic autosomal fragments (>500,000 base pairs) for further analyses. The program MEDICC was used (with default settings) to infer the evolutionary hierarchy from the CNA profiles; we then annotated each branch (i.e., monophyletic clade) of the resulting evolutionary hierarchy with a confidence score (referred as subsampling confidence) using an iterative chromosome subsampling approach. For this approach, the steps described above were iterated 30 times using random subsamples (without replacement) of 15 autosomes at the time; the frequency at which a selected monophyletic clade in the source evolutionary hierarchy appeared in the sampling simulations indicated the confidence of that clade. The branches of the source evolutionary hierarchy with a low confidence score (<0.5; i.e., majority rule) were then collapsed to their most proximal predicted ancestral cell population (referred from now on as the ancestor). The majority of the branches derived from the predicted evolutionary hierarchies (Fig. 2) had a subsampling confidence higher than 0.5 (i.e., appeared in greater than 50% of the sampling simulations). The only exceptions were the branching nodes L, H and v from patient 167 that had a subsampling confidence of 0.46 and were collapsed to node iv, and for the branching nodes B and D from patient 299 that had a subsampling confidence of 0.43 and were collapsed to node i. The shoulder metastasis from patient 299 was placed in the evolutionary hierarchy based on additional genomic information from Hong et al. Such an approach was employed because this particular sample was analysed with a higher density SNP array, compared with the other samples in the same evolutionary hierarchy, introducing a clear bias that prevented high confidence predictions, based solely on SNP array information.

**Comparative analyses of copy number profiles.** For identifying the potential driver CNAs across transitions between histo-pathological categories, we selected for each evolutionary hierarchy the representative ancestral population for each category; with the criteria that it should originate all (or the majority) of sampled clones belonging to such category. For each transition between histo-pathological categories, we obtained the increase/ decrease CNA profile subtracting two consecutive representatives (e.g., central subtracted from extraprostatic extension). Then, we consistently fragmented the resulting increase/decrease CNA profiles to allow direct comparison and tested each genomic fragment for differences across histo-pathological categories using analysis of variance (ANOVA; function aov from R platform). All changes with a false-discovery corrected (method Benjamini, Hochberg, and Yekutieli p-value <0.25) are reported in Table 2.

**Validation.** We independently validated the CNA profiles inferred using ASCAT from SNP array data using the copy numbers predicted from Multiplex Ligation-dependent Probe Amplification (MLPA) technology on 58 genomic regions spanning 39 genes (kit ID P294- B1) over the samples 167C, 167D, 167E, 167G, 167H, 167I, 167L, 179C, 179D, 179E, 179G, 179L, 421C, 421E, 421F, 421G, 421H and 421L. MLPA was performed as per vendor’s protocol (MRC-Holland, Amsterdam, The Netherlands). We normalized the raw MLPA intensities for probe length using the function mlpaNorm (using the method “slope correction”) from the R package MLPAstats, and calculated the ratio between the normalized MLPA intensity values for genes in tumour samples and benign, control samples (i.e., 167B, 179B, 421A). Then, from those ratios, we inferred the gene CNs using thresholds adjusted for tumour ploidy (Fig. S3) according to the formula ratio = (ploidy-2)/2. Each branch of the
evolutionary hierarchies was considered validated if a significant proportion of genes in a variation in CN was validated; a gene was considered validated if the copy number obtained using MLPA and SNParray was the same. Evolutionary hierarchies for the patients 001, 177, 299 and 498 were indirectly validated comparing their topologies with the topologies of the evolutionary hierarchies obtained with somatic point mutations and structural variation analysis identified from whole genome sequencing.11

Data and computational algorithms. The raw data of the SNP arrays for patients 167, 179, 421 can be retrieved at ega-archive.org with the code EGAS00001001801. Theinformatics code used for the analyses in this work can be retrieved at github.com/stemangiola/PC-lymph-node-mets-evolution.

References
1. Paget, S. The Distribution Of Secondary Growths In Cancer Of The Breast. The Lancet 133, 571–573 (1889).
2. Briganti, A. et al. Updated nomogram predicting lymph node invasion in patients with prostate cancer undergoing extended pelvic lymph node dissection: the essential importance of percentage of positive cores. Eur Urol 61, 480–487, doi: 10.1016/j.euro.2011.10.044 (2012).
3. Briganti, A. et al. Lymphatic spread of nodal metastases in high-risk prostate cancer: The ascending pathway from the pelvis to the retroperitoneum. Prostate 72, 186–192, doi: 10.1002/pros.21420 (2012).
4. Moschini, M. et al. Natural History of Clinical Recurrence Patterns of Lymph Node-Positive Prostate Cancer After Radical Prostatectomy. Eur Urol 69, 135–142, doi: 10.1016/j.eururo.2015.03.036 (2016).
5. James, N. D. et al. Survival with Newly Diagnosed Metastatic Prostate Cancer in the "Docetaxel Era": Data from 917 Patients in the Control Arm of the STAMPEDE Trial (MRC PR08, CRUK/06/019). Eur Urol 67, 1028–1038, doi: 10.1016/j.eururo.2014.09.032 (2015).
6. Messing, E. M. et al. Immediate versus deferred androgen deprivation treatment in patients with node-positive prostate cancer after radical prostatectomy and pelvic lymphadenectomy. Lancet Oncol 7, 472–479, doi: 10.1016/S1470-2045(06)70700-8 (2006).
7. Halsted, W. S. I. The Results of Operations for the Cure of Cancer of the Breast Performed at the Johns Hopkins Hospital from June, 1889, to January, 1894. Ann Surg 20, 497–555 (1894).
8. Schiavina, R. & Brunocilla, E. Prostate cancer: lymph node metastases: not always the same prognosis. Nat Rev Urol 10, 435–436, doi: 10.1038/nrurol.2013.127 (2013).
9. Datta, K., Muders, M., Zhang, H. & Tindall, D. J. Mechanism of lymph node metastasis in prostate cancer. Future Oncol 6, 823–836, doi: 10.2217/fon.10.33 (2010).
10. Gundem, G. et al. The evolutionary history of lethal metastatic prostate cancer. Nature 520, 353–357, doi: 10.1038/nature14347 (2015).
11. Hong, M. K. et al. Tracking the origins and drivers of subclonal metastatic expansion in prostate cancer. Nat Commun 6, 6605, doi: 10.1038/ncomms7605 (2015).
12. Nowell, P. C. The clonal evolution of tumor cell populations. Science 194, 23–28 (1976).
13. Bielas, J. H., Loeb, K. R., Rubin, B. P., True, L. D. & Loeb, L. A. Human cancers express a mutator phenotype. Proc Natl Acad Sci U S A 103, 18238–18242, doi: 10.1073/pnas.0607057103 (2006).
14. Heng, H. H. et al. Stochastic cancer progression driven by non-clonal chromosome aberrations. J Cell Physiol 208, 461–472, doi: 10.1002/jcp.20685 (2006).
15. Reiter, J. G. et al. Reconstructing phylogenies of metastatic cancers. bioRxiv, doi: 10.1101/048157 (2016).
16. Naxerova, K. & Jain, R. K. Using tumour phylogenetics to identify the roots of metastasis in humans. Nat Rev Clin Oncol 12, 258–272, doi: 10.1038/nrclinonc.2014.238 (2015).
17. Hong, W. S., Shpak, M. & Townsend, J. P. Inferring the Origin of Metastases from Cancer Phylogenies. Cancer Res 75, 4021–4025, doi: 10.1158/0008-5472.CAN-15-1889 (2015).
18. Schwarz, R. F. et al. Phylogenetic quantification of intra-tumour heterogeneity. PLoS Comput Biol 10, e1003355, doi: 10.1371/journal.pcbi.1003355 (2014).
19. Sato, K. et al. Clinical significance of alterations of chromosome 8 in high-grade, advanced, nonmetastatic prostate carcinoma. J Natl Cancer Inst 91, 1574–1580 (1999).
20. Williams, J. L., Greer, P. A. & Squire, J. A. Recurrent copy number alterations in prostate cancer: an in silico meta-analysis of publicly available genomic data. Cancer Genet 207, 474–488, doi: 10.1016/j.cancergen.2014.09.003 (2014).
21. Nickerson, M. L. et al. Somatic alterations contributing to metastasis of a castration-resistant prostate cancer. Hum Mutat 34, 1231–1241, doi: 10.1002/humu.22346 (2013).
22. Takayama, K. et al. TET2 repression by androgen hormone regulates global hydroxymethylation status and prostate cancer progression. Nat Commun 6, 8219, doi: 10.1038/ncomms9219 (2015).
23. Pelenaris, S., Khan, M. & Evans, G. c-MYC: more than just a marker of life and death. Nat Rev Cancer 2, 764–776, doi: 10.1038/nrc7904 (2002).
24. Santra, M. K., Waiyapee, N. & Green, M. R. F-box protein FBXO31 mediates cyclin D1 degradation to induce G1 arrest after DNA damage. Nature 459, 722–725, doi: 10.1038/nature08011 (2009).
25. Xiao, X., Li, B. X., Mitton, B., Ikeda, A. & Sakamoto, K. M. Targeting CREB for cancer therapy: friend or foe. Curr Cancer Drug Targets 10, 384–391 (2010).
26. Massoumi, R. CYLD: a deubiquitination enzyme with multiple roles in cancer. Future Oncol 7, 285–297, doi: 10.2217/fon.10.187 (2011).
27. Kikuchi, M. et al. TRIM24 mediates ligand-dependent activation of androgen receptor and is repressed by a bromodomain-containing protein, BRD7. In prostate cancer cells. Biochim Biophys Acta 1793, 1828–1836, doi: 10.1016/j.bbamcr.2009.11.001 (2009).
28. Osunkoya, A. O., Adsay, N. V., Cohen, C., Epstein, J. I. & Smith, S. L. MUC2 expression in primary mucinous and nonmucinous adenocarcinoma of the prostate: an analysis of 50 cases on radical prostatectomy. Mod Pathol 21, 789–794, doi: 10.1038/modpathol.2008.47 (2008).
29. Brastianos, P. K. et al. Genomic Characterization of Brain Metastases Reveals Branched Evolution and Potential Therapeutic Targets. Cancer Discov 5, 1164–1177, doi: 10.1158/2159-8290.CD-15-0369 (2015).
30. Furuya, Y., Akakura, K., Akimoto, S. & Ito, H. Prognosis of patients with prostate carcinoma presenting as nonregional lymph node metastases. Urol Int 61, 17–21 (1998).
31. Gandaglia, G. et al. Impact of the Site of Metastases on Survival in Patients with Metastatic Prostate Cancer. Eur Urol 68, 325–334, doi: 10.1016/j.eururo.2014.07.020 (2015).
32. Barekati, Z. et al. Methylation signature of lymph node metastases in breast cancer patients. BMC Cancer 12, 244, doi: 10.1186/1471-2407-12-244 (2012).
33. Senkus, E. et al. Primary breast cancer: ESMO Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 24 Suppl 6, vi7–23, doi: 10.1093/annonc/mdt284 (2013).
34. Urrutia, G. et al. Epigenetic variations in breast cancer progression to lymph node metastasis. Clin Exp Metastasis 32, 99–110, doi:10.1007/s10585-015-9695-4 (2015).
35. Budaus, L. et al. Initial Experience of Ga-PSMA PET/CT Imaging in High-risk Prostate Cancer Patients Prior to Radical Prostatectomy. Eur Urol, doi: 10.1016/j.eururo.2015.06.010 (2015).
36. Maurer, T. et al. Diagnostic Efficacy of Gallium-PSMA-PET compared to Conventional Imaging in Lymph Node Staging of of 130 consecutive Patients with Intermediate to High-Risk Prostate Cancer. J Urol, doi: 10.1016/j.juro.2015.12.025 (2015).
37. Rowe, S. P. et al. PSMA-Based Detection of Prostate Cancer Bone Lesions With (18F)-DCFPyL PET/CT: A Sensitive Alternative to (99m)Te-MDP Bone Scan and Na(18)F PET/CT? Clin Genitourin Cancer 14, e115–e118, doi: 10.1016/j.clgc.2015.09.011 (2016).
38. Sterzing, F. et al. (68)Ga-PSMA-11 PET/CT: a new technique with high potential for the radiotherapeutic management of prostate cancer patients. Eur J Nucl Med Mol Imaging 43, 34–41, doi: 10.1007/s00259-015-3188-1 (2016).
39. Verburg, F. A. et al. Extent of disease in recurrent prostate cancer determined by [Ga]PSMA-HBED-CC PET/CT in relation to PSA levels, PSA doubling time and Gleason score. Eur J Nucl Med Mol Imaging, doi: 10.1007/s00259-015-3240-1 (2015).
40. Zamboglou, C. et al. MRI versus Ga-PSMA PET/CT for gross tumour volume delineation in radiation treatment planning of primary prostate cancer. Eur J Nucl Med Mol Imaging, doi: 10.1007/s00259-015-3257-5 (2015).
41. Hong, M. K. et al. Error rates in a clinical data repository: lessons from the transition to electronic data transfer – a descriptive study. BMJ Open 3, doi: 10.1136/bmjopen-2012-002406 (2013).
42. Ao, M. et al. Cross-talk between paracrine-acting cytokine and chemokine pathways promotes malignancy in benign human prostatic epithelium. Cancer Res 67, 4244–4253, doi: 10.1158/0008-5472.CAN-06-3946 (2007).
43. Van Loo, P. et al. Allele-specific copy number analysis of tumors. Proc Natl Acad Sci USA 107, 16910–16915, doi: 10.1073/pnas.1009843107 (2010).
44. Benjamini, Y. & Hochberg, Y. Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. Journal of the Royal Statistical Society. Series B (Methodological) 57, 289–300 (1995).
45. Van Opstal, D. et al. Multiplex ligation dependent probe amplification (MLPA) for rapid distinction between unique sequence positive and negative marker chromosomes in prenatal diagnosis. Mol Cytogenet 4, 2, doi: 10.1186/1755-8166-4-2 (2011).
46. Caceres, A., Armengol, L., Villatoro, S. & Gonzalez, J. R. MLPAstats: an R GUI package for the integrated analysis of copy number alterations using MLPA data. BMC Bioinformatics 12, 147, doi: 10.1186/1471-2105-12-147 (2011).

Acknowledgements
We thank M. Kerger and P. Xibberas for tumour sample collection and patient disease annotation. We thank all members of the CMH laboratory for helpful comments and editing of this manuscript. This work was supported, in part, by NHMRC grants 1047581 and 1104010 to CMH and 1024081 to NMC. The Ovarian Cancer Action [BritROC project grant: G.M]; The University of Cambridge, Cancer Research UK, Hutchinson Whampoa Limited, CRUK core grant C14303/A17197 (CRUK CI institute core grant, GM).

Author Contributions
S.M., M.K.H.H., N.M.C., G.M. and C.M.H. designed studies and conducted experiments. M.K.H.H conducted experiments and analysed patient records. N.K. performed MLPA experiments. S.M., M.C., A.R., A.C. and G.M. analysed data and reviewed the paper. S.M., G.M., N.M.C and C.M.H. wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep
Competing financial interests: The authors declare no competing financial interests.
How to cite this article: Mangiola, S. et al. Comparing nodal versus bony metastatic spread using tumour phylogenies. Sci. Rep. 6, 33918; doi: 10.1038/srep33918 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/}
© The Author(s) 2016
Minerva Access is the Institutional Repository of The University of Melbourne

Author/s:
Mangiola, S; Hong, MKH; Cmero, M; Kurganovs, N; Ryan, A; Costello, AJ; Corcoran, NM; Macintyre, G; Hovens, CM

Title:
Comparing nodal versus bony metastatic spread using tumour phylogenies

Date:
2016-09-22

Citation:
Mangiola, S., Hong, M. K. H., Cmero, M., Kurganovs, N., Ryan, A., Costello, A. J., Corcoran, N. M., Macintyre, G. & Hovens, C. M. (2016). Comparing nodal versus bony metastatic spread using tumour phylogenies. SCIENTIFIC REPORTS, 6 (1), https://doi.org/10.1038/srep33918.

Persistent Link:
http://hdl.handle.net/11343/118616

License:
CC BY