

A case of raised PSA with ALP

HOSP #	Lab no. SA04016354	WARD	Orthopaedic Clinic
CONSULTANT	Jody Rusch	DOB/AGE	61y Male

Abnormal Result

PSA: 846.5 ug/L

ALP: 284 U/L (53 – 128)

Presenting Complaint

Painful "lumps" in groin + constipation

Spine pain

History

Smoker (>45 years)

No other co-morbidities

6/12 history of generalized body pain (mostly spine)

Red Flags (weightloss, night pain not responding to analgesia)

Examination

O/E: Pallor (Hb 8.6), Wasted. Clinically painful bilateral inguinal lymph nodes PR: normal tone, no masses, no blood, prostate smooth

Laboratory Investigations

Na	138 mM
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K	4,7 mM
Cl	101 mM
Urea	10,3 mM
Creat	69 uM
eGFR by MDRD	>60 ml/min/m2
eGFR by CKDEPI	97 ml/min/m2
Ca	2,26 mmol/L
Mg	1,03 mmol/L
Phos	1,01 mmol/L
Total prot	73 g/L
Alb	37 g/L
Total bili	3 umol/L
Conj bili	2 umol/L
ALT	15 U/L (10-40)
AST	19 U/L (15-40)
ALP	284 U/L (53 – 128)
GGT	76 U/L (<68)
LD	345 U/L
CRP	52 mg/L (<10)
Total PSA	846.5 ug/L (<4)
TSH	1,33 mIU/L (0.27 – 4.2)
Hb	5.6 g/dL
MCV	88.3 fL
WCC	7.57 cells/uL

Table 1 – Blood results on 06/07/2020

Other Investigations

Chest X-Ray: Left hilar opacities

X-ray of the limbs: Global lytic lesions involving both

proximal femurs



Figure 1 – Lytic lesion seen in the centre of the thoracic vertebral body.

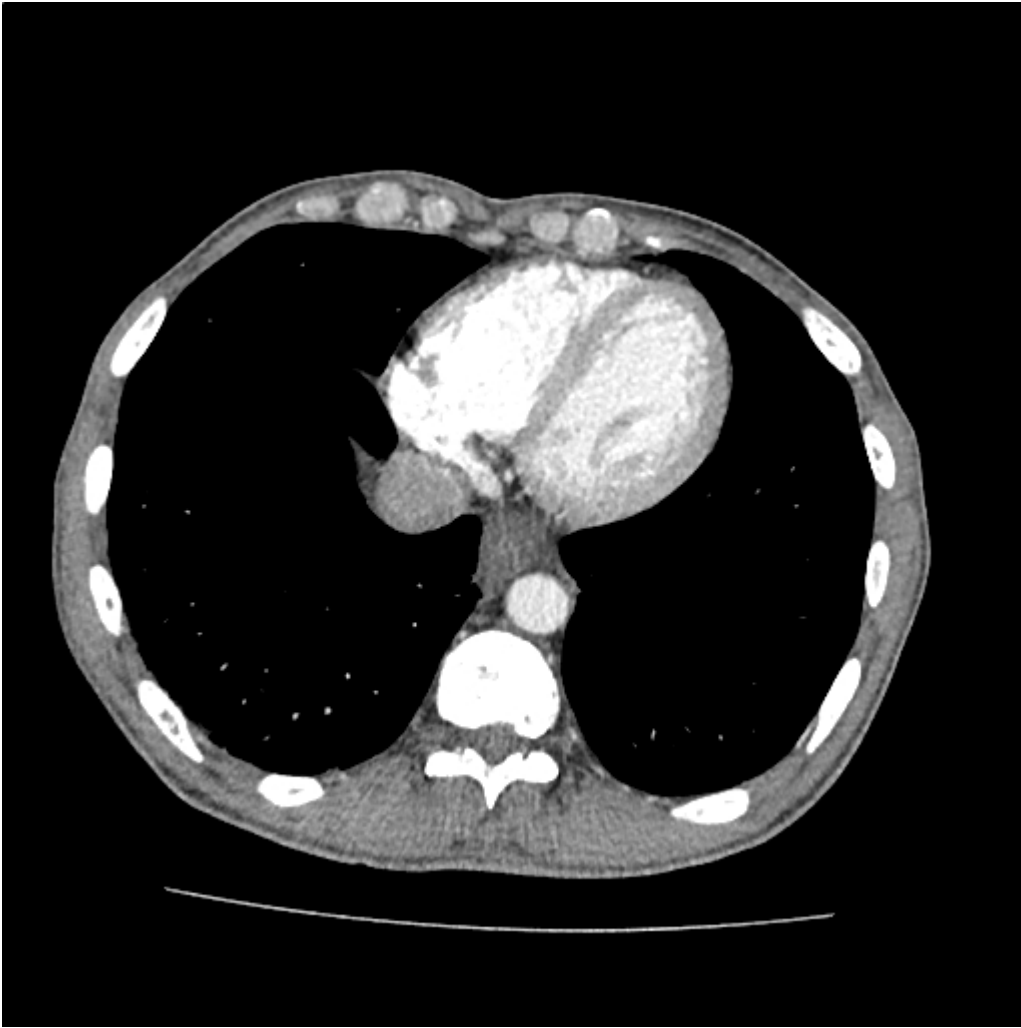


Figure 2 – Included for comparison with Figure 1 – not as big lytic lesion seen.

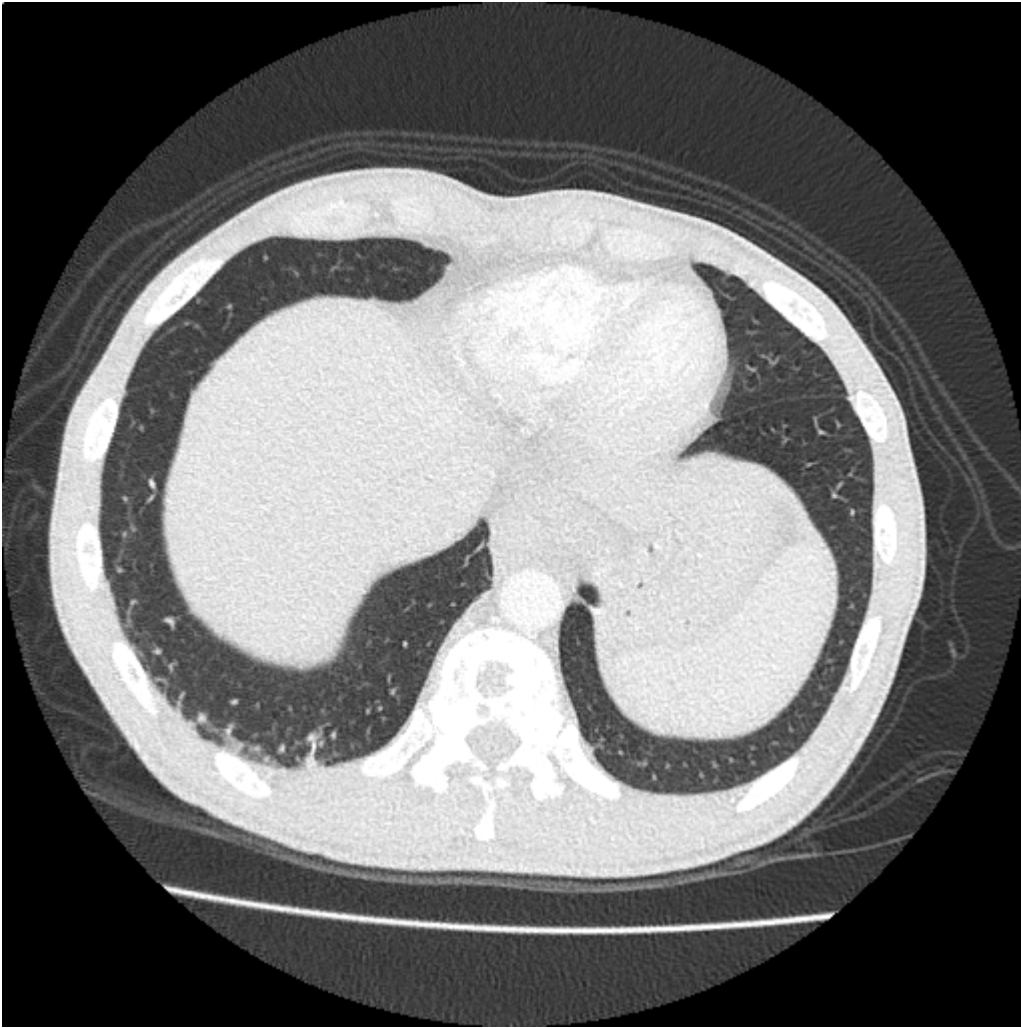


Figure 3 – MRI image of the same thoracic vertebral body as shown in Figure 1.

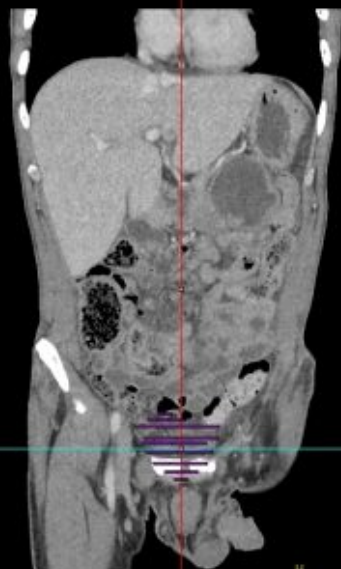
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09-Jul-2020

FLP



W 400

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W 400

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ID	Type	Value
M1	Volume measurement	62.40 cm ³

Figure 4 – Transverse and coronal views of the CT scan with the outline of the prostate marked in yellow (left middle) and purple lines (right top and bottom)



Figure 5 – Small lytic lesions visible in the proximal femur.

Prostate biopsy

- **MACROSCOPY:** Specimen consists of two cores, the longest measuring 12mm in length.
- **MICROSCOPY** Sections show 2 prostatic cores, both infiltrated by a prostatic adenocarcinoma.
- % Ca core 1: 90%
- % Ca core 2: 60%
- Gleason score: 5 + 4
- Grade group: 5
- High grade PIN: Not seen
- Seminal vesicle: Not seen
- Perineural invasion: Present
- Fat (extraprostatic) involvement: Not seen
- **PATHOLOGICAL DIAGNOSIS:**
 - **Prostate, needle biopsy: Prostatic acinar adenocarcinoma**

Final Diagnosis

Metastatic Prostate Carcinoma with multiple metastases to the

bones (thoracic spine and both femurs).

Take Home Message

Prostate-specific antigen (PSA, also known as kallikrein III, seminin, semenogelase, γ -seminoprotein and P-30 antigen) is a 34-kD glycoprotein produced almost exclusively by the prostate gland. It is a serine protease enzyme.

Most PSA in the blood is bound to serum proteins. A small amount is not protein-bound and is called 'free PSA'. In men with prostate cancer, the ratio of free (unbound) PSA to total PSA is decreased. The risk of cancer increases if the free to total ratio is less than 25%.

The lower the ratio is, the greater the probability of prostate cancer. Measuring the ratio of free to total PSA appears to be particularly promising for eliminating unnecessary biopsies in men with PSA levels between 4 and 10 mg/L.

ALP (alkaline phosphatase) is well known to be a marker of ductal hepatic damage. ALP, being an isozyme, however has its origin from various tissue sources in the body. It is present in the liver, bile duct, kidney, bone, intestinal mucosa and placenta. The majority of ALP in serum is from either skeletal or liver origin. In adults the major form is from liver and in children the major form is from the skeleton.

Blood levels of alkaline phosphatase increase by two to four times during pregnancy. This is a result of additional alkaline phosphatase produced by the placenta.

If it is unclear why alkaline phosphatase is elevated, isoenzyme studies using electrophoresis can confirm the source of the ALP. It would likely in this patient be quite clear that the raised ALP would be due to the excess

leakage from the osteolytic lesions from the metastases, but who knows, the patient may have had a beer or five in the preceding 3 weeks leading up to the bloods being drawn. The fact that the other liver enzymes are near-normal, makes alcohol consumption less likely though.