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CLINICAL SIGNIFICANCE OF HIGH GRADE AND LOW GRADE PROSTATE INTRAEPITHELIAL NEOPLASIA

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Abstract. Clinical significance of high grade and low grade prostate intraepithelial neoplasia. Melnychuk M.P.
Such premalignant conditions of prostate cancer (PC) as prostate intraepithelial neoplasia (PIN) are classified between benign and malignant ones. Contemporary evidence wheather PIN develops malignancy is limited and (LGPIN) data present varied results. Morphological and clinical differencies between high (HGPIN) and low grade PIN specimens in the prostate remain unclear. Aim of the work – to determine clinical significance and progression ability of high grade and low grade prostate intraepithelial neoplasia. The results of examination of 276 patients with PIN (152 patients with high grade PIN and 134 patients with low grade PIN) were assessed comparatively. During a 3 year follow-up repeated prostate biopsies were performed with 6 months interval to detect PC. Initial and repeated multifocal transrectal prostate biopsies from 12 samples were performed under transrectal ultrasonic guidance. There were statistically
Prostate cancer (PC) occupies the second place in the structure of malignancies (after lung cancer) in men worldwide, counting 1,276,106 new cases and causing 358,989 deaths (3.8% of all deaths caused by cancer in men) in 2018 [7].

Prostate intraepithelial neoplasia (PIN) which was described in 1969 [1] is a preneoplastic proliferation process of prostatic epithelium that is bounded to prostatic ducts or acini (glands). According to degree of pathological cellular changes, PIN is classified as low grade PIN (LGPIN) and high grade PIN (HGPIN). Despite numerous investigations the role of PIN in prostate cancerogenesis is being strongly discussed. There is abundant data supporting the statement that high-grade PIN (HGPIN) is a precursor lesion to adenocarcinoma of the prostate [9]. According to investigation data, HGPIN and PC have similar epidemiological and clinical features [16]. Like PC, the incidence of HGPIN depends on age and is higher in African-American men. HGPIN is more frequently detected in postoperative prostates along with PC than without it [18]. Also HGPIN is often multifocal and is located in the peripheral lateral part of the prostate gland [14]. HGPIN has similar to PC molecular and genetic changes that were investigated by researchers [5].

The incidence of high grade and low grade PIN is still under investigation. Despite the importance of recognizing PIN, there is still insufficient standardization in diagnostics methods and their description. There is large variability in reports of different pathologists due to subjectivity of diagnosis of PIN [13]. This variance depends on few peculiarities, such as type of specimen, preparation of tissue, selection of patients for biopsy [19]. Interobserver variability makes problematic the investigation of epidemiology and clinical diagnosis of prostate adenocarcinoma [10]. Tan et al reviewing “benign” needle core biopsies specimens demonstrated that 75% of HGPIN lesions were initially missed by reporting pathologists [2]. Moreover, Krones et al discovered a 30-patient oversight of HGPIN out of 3251 prostate biopsies. According to data of researchers HGPIN formed 34.5% of the missed cancer which is detected on subsequent biopsies. According to data of researchers HGPIN formed 34.5% of the missed lesions identified in the study [6]. Because of lack of standardization in case of low-grade PIN, urologists do not routinely report this precancerous state, only for investigation aims [11].

Clinical significance of PIN depends on its malignant potential and on its association with cancer which is detected on subsequent biopsies. Analysis of scientific works that were conducted before mid-1990s and assessed cancer risk on repeated prostate biopsies after diagnosis of isolated HGPIN demonstrated a great data variety – from 27% to 100%. After the prevalent introduction of extended core biopsy techniques, it has been established that there is a lower risk of PC detection on repeated biopsies [1]. Lower incidence of prostate adenocarcinoma detection on repeated biopsies was connected with prostate screening programs using widespread prostate-specific antigen (PSA) and as a
result in insignificant, small-volume cancer at
diagnosis, and introduction of extended biopsy
techniques with investigation of the peripheral
lateral zone of the prostate gland [17].

The spread of PIN in prostate gland is assessed as
a risk factor of malignization. To the opinion of
Merrimen JL et al. the number of positive cores that
contain HGPIN is the one pathological factor that
predicts a higher risk of PC on repeated biopsy. The
presence of multifocal high-grade PIN on prostate
biopsy (involving 2 or more cores) represents a risk
factor for detection of cancer on repeated biopsies [17].
HGPIN with 3 cores or more is associated with a
sufficiently high risk of cancer and requires re-biopsy
within a year of the initial PIN diagnosis. For cases
with one or two cores of HGPIN on needle biopsy, it is
recommended that men should not have a obligatory
repeated needle biopsy during the first year after the
diagnosis of HGPIN, in case of absence of other
clinical and morphological risk factors of PC [12].

Although HGPIN is linked to PC, its malignant
potential has been heavily debated. In fact, some
studies discuss the fact of association of isolated
HGPIN with a high risk of prostate adenocarcinoma on
repeated biopsies [5, 10, 11]. Contemporary evidence
regarding the etiology, natural history, surveillance,
and management of PIN is limited. There is an
abundance of compelling evidence to suggest that
HGPIN is a precursor of PC. More studies with long
follow-up period are needed to establish the role of
high and low grade PIN in cancerogenesis of prostate.

The aim of the study was to investigate malignant
transformation rates and clinical significance of high
grade and low grade prostate intraepithelial neplasia.

**MATERIALS AND METHODS OF RESEARCH**

We studied 276 patients aged 48 to 76 years
(mean age 65.6±1.4 years) with PIN. All patients
were divided into two groups according to PIN
grade. The first group consisted of 152 patients with
high grade PIN. The second research group
consisted of 134 patients with low grade PIN.
Diagnosis was confirmed morphologically after
transrectal multifocal prostate biopsy that was
performed according to PC suspicion. The method-
dics of prostate biopsy and morphological diagnosis
were established according to European Association
of Urology (EAU) Guidelines and Transrectal
Ultrasound Guided Biopsy of the Prostate [3, 4]
Within a 3 year follow-up there were performed
prostate rebiopsies with 6 months’ interval. The aim
of rebiopsies was to assess morphological changes in
prostate tissue and to detect PC. Initial and repeated
transrectal prostate biopsies were performed under
transrectal ultrasonic guidance with 12 samples.

Statistical analysis of the results was performed
using “Statistica” (version 6.0, Statsoft Inc, USA.
Licence AXXR712D833214FAN5) software. The
significance of received results between groups was
evaluated by Mann– Whitney test, p<0.05 was
considered as statistically significant [15].

**RESULTS AND DISCUSSION**

During a 3-years’ follow-up in 75 (27.2%) of
276 patients with PIN in repeated biopsies, PC
was detected. The aim was to determine
correlation between grade of PIN and rate of
malignant transformation. PC was diagnosed in 64
(42.1%) patients with HGPIN and in 11 (8.2%)
patients with LGPIN (Fig. 1).

![Graph](image-url)
To study correlation between malignization rate and spread of PIN lesions in prostate tissue all patients were divided into groups by number of PIN positive cores. 106 patients composed the first group who had 1-2 PIN samples, the second group (93 patients) – 3-4 samples, the third group (77 patients) – 5 and more samples. Analysis showed that PC was detected in 16 (21.3%) of patients with 1-2 PIN lesions, in 23 (30.7%) of patients with 3-4 PIN lesions and in 36 (48%) of patients with 5 and more PIN lesions (Fig. 2).

Thus analysis of PIN spread in prostate tissue that was determined as a number of PIN positive biopsy samples established direct correlation between PIN spread and subsequent PC. Most frequently malignant transformation was observed in patients who had 5 and more PIN samples initially. In group with 3-4 PIN samples PC, was detected by 22.1% and in group with 1-2 PIN samples – by 31.7 less frequently.

The follow-up period was 3 years. The aim was to study the terms of high and low grade PIN malignant transformation and adenocarcinoma diagnostics. It was found that among 64 PC cases in patients with HGPIN that were detected during 3-year follow-up period in 28 (43.8%) patients adenocarcinoma was diagnosed during the period from 12 months to 18 months, in 16 (25%) patients from 6 to 12 months, in 11 (17.2%) patients from 18 to 24 months, in 4 (6.3%) from 24 to 30 months and from 30 to 36 months and in 1 (1.6%) PC was diagnosed in 6 months after HGPIN diagnose.

Among 11 PC cases in patients with LGPIN that were detected during 3-year follow-up period in 5 (45.5%) patients adenocarcinoma was diagnosed during the period from 24 to 30 months, in 3 (27.3%) – from 30 to 36 months, in 2 (18.2%) patients – from 18 to 24 months, in 1 (9.1%) patients – from 12 to 18 months. During the first year after initial biopsy in patients with LGPIN, PC was not detected.

Thus obtained data evidence that the terms of PIN transformation into adenocarcinoma variate from 1 to 3 years. The difference was observed between median terms of HGPIN and LGPIN malignant transformation. In patients with HGPIN PC was diagnosed mainly during the period from 12 to 24 months (the second year follow-up), namely 39 (60.9%) PC cases. Conversely, in patients with LGPIN the main amount of adenocarcinoma was detected later – during the period from 24 to 30 months, namely 5 (45.5%) PC cases.

Clinical significance of precancerous disease is determined by its malignant transformation ability. Numerous investigations that were performed to make clear PIN malignization rate came to opposite conclusions because of different research methods, namely histological and statistical differences. Historically it is difficult to compare morphological data because a lot of researchers used different approaches to describe such premalignant states as high and low grade PIN, ASAP, intraductal carcinoma. Most studies were analyzed with insufficient material for reliable statistical assessment including data of prostate biopsies, transurethral prostate resection, autopsies.
In our study we tried to follow last recommendations and classifications of premalignant prostate states and to use appropriate statistical methods. It was determined that clinical significance of PIN correlates with its grade. HGPIN is a precancerous pathology with malignization rate of 42.1% while follow-up of patients with LGPIN demonstrated insignificant malignization ability of 8.2%. Obtained data correlate with those of De Marzo et al. [16]. Besides it was established that PC rate in patients with HGPIN depends on spread of PIN in prostate, namely on the number of positive samples. The highest risk of PC diagnose is in case of 5 and more PIN samples. Our data correlate with research work of Zhou M. et al. [8].

The terms of malignization differ in patients with high and low grade PIN. Our study showed that cancerogenesis is a gradual process that proceeds through sequenced stages from benign tissue through LGPIN to HGPIN and finally to prostate adenocarcinoma.

It is important to distinguish a group of high malignization risk among patients with PIN based on immunohistochemical investigation, PSA, MRI data. These patients need detailed precise examination and treatment to prevent malignant transformation in every individual case.

CONCLUSIONS
1. High grade prostate intraepithelial neoplasia is a prostate cancer precursor with high progression ability. Malignization rate in patients with high grade prostate intraepithelial neoplasia amounted 42.1% during a 3-year follow-up and was by 33.9% higher than in low grade prostate intraepithelial neoplasia patients.
2. Low grade prostate intraepithelial neoplasia should be considered an initial cancerogenesis process with mainly benign tissue features. Malignization rate in patients with low grade prostate intraepithelial neoplasia amounted 8.2% during a 3-year follow-up.
3. The spread (extent) of high grade prostate intraepithelial neoplasia lesions within prostate gland is a malignization risk factor. It was established a direct correlation between with prostate cancer rate and number of biopsy cores high grade prostate intraepithelial neoplasia positive samples prostate gland.
4. Low and high grade prostate intraepithelial neoplasia are gradual stages of cancerogenesis. The mean malignization term of high grade prostate intraepithelial neoplasia is 18 months and of low grade prostate intraepithelial neoplasia – 30 months.
5. Prostate intraepithelial neoplasia grade determines its clinical significance. While low grade prostate intraepithelial neoplasia has low malignization potential, high grade prostate intraepithelial neoplasia possesses morphological and clinical characteristics of adenocarcinima.

Conflict of interests. The authors declare no conflict of interest.
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СПИСОК ЛІТЕРАТУРИ

5. Extent of baseline prostate atrophy is associated with lower incidence of low and high-grade prostate cancer.


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