

**Sub: Inviting objections, comments for Procurement of Anti- Plagiarism Software.**

The Institute is in process to purchase **Anti- Plagiarism Software and Plasma/Low Hb Analyzer** for the Institute.

S. No.	Platform	PAC Certification by
1.	Anti- Plagiarism Software	Turnitin, LLC
2.	Plasma/Low Hb Analyzer	Hemocue India

The PAC Certifications by Company and as well as user are attached.

The above documents with specifications are being uploaded for open information and also to submit online objections, comments, if any from any manufacturer regarding proprietary nature of **Anti- Plagiarism Software and Plasma/Low Hb Analyzer** giving Advt. reference No. KSSSCI/Tender-05/2025-26. The comments should be received in the name of the Director, KSSSCI, Sultanpur Road, Lucknow-226002 on or before **05.06.2025 up to 04:00 pm**, failing which it will be presumed that any other vendor is having no comments, objections for above purchase on proprietary basis and case will be decided on merits.

- 1-Specification of items mentioned above.
- 2- PAC Certificate.

## **Specification of Anti-Plagiarism software**

The software/ Cloud based software should be able to the check for plagiarized content and should support the following minimum requirement:-

### **(A) Essential Features of Software:**

1. It should be a web-based interface hosted by a secure data center.
2. It should be accessible from any place through login/password control
3. It should be capable of searching all medical literature available on the day of search on the internet in the form of scholarly journals, periodicals, books, reports, and e-books with valid DOI even if it is password controlled.
4. It should be capable of loading, checking, and assessing large file sizes in different formats such as Microsoft Word, HTML, XML, rich text format, PDF, Open Office, etc.
5. It should have a facility for plagiarism checks for commonly used file formats such as Word, PDF, Text files, Web pages, Open Office documents, etc.
6. It should have a word-to-word matching ability with the exclusion of common words, phrases, and medical terms. It should be able to identify and report copy-paste with source identification.
7. It should provide an overlap/similarity percentage with a detailed report of the sources that are similar to the article being assessed. This report should be download-able.
8. There should not be any word limit for plagiarism check in one go.
9. The company should have online technical assistance available 24X7 for all technical issues.
10. The company should have developed or provided similar software for other institutes of repute like AIIMS Delhi, PGI Chandigarh, CMC Vellore, JIPMER etc. and preferably international health institutes of repute like NIH & Welcome Trust.
11. The software should include all types of available medical literature publisher's databases listed in the Cross Reference & ProQuest websites in their similarity check database coverage of medical literature databases.
12. Software should have advanced AI content detection capabilities.

*[Signature]*

*[Signature]*

**(B) Features of the report generated:**

1. Originally report generation with date & time, name, location, word count, and author of the document that has been checked.
2. A summary of the relative amount of plagiarized portion of the checked document, the original document, and the referenced part if any should be shown in different colors/ highlighted portions in the source document.
3. The report should show the original and copy source in side-by-side columns.
4. There should not be any change in the format after the report is generated.
5. The software should be demonstrated to concerned authority, Library Committee, and A proper training schedule is required for Library staff as per the requirement.

**(C) End users access :**

1. Subscription period will be 12 months from the date of service activation.
2. The Company should provide access of at least 5,000 submissions during the subscription period.

The vendor must provide options for extra submissions, if required, during the subscription period.

**(D)** Free trial access to software for a limited period to be provided for technical evaluation.

**(E)** The vendor must provide Strongest DATA security and DATA privacy of the industry and compliance to global standards like COPPA, FERPA, GDPR , SSL encryption, AICPA, SOC etc.

**(F)** The vendor must provide publications a collection of over 80 million articles from library databases, textbook publishers, digital reference collections, subscription-based publications, homework helper sites, and books.

*[Signature]*

*[Signature]*

(4)

## Kalyan Singh Super Specialty Cancer Institute

कल्याण सिंह अति विशिष्ट कैंसर संस्थान

C.G. City, Sultanpur Road, Lucknow-226002

सी०जी० सिटी, सुल्तानपुर रोड, लखनऊ- 226002

(An Autonomous Institute of the Govt. of Uttar Pradesh)

(उत्तर प्रदेश सरकार का स्वायत्तशासी संस्थान)

Email: jdmm.sscih@gmail.com

### Proprietary Article Certificate (PAC) for Items/Goods

(1) The indented goods are manufactured by  
M/s. TURJINTIN, LLC.

(2) No other make or model is acceptable for the following reasons:

- a) Unparalleled database of Internet and scholarly publishers.
- b) Proprietary web crawler and private repository.
- c) Data security (FERPA & COPPA compliant).

Signature  
30-MAR-2018  
.....  
(Signature of Indentor)

.....  
.....  
(Signature of HOD)



## Additional term and conditions for Anti-Plagiarism software

1. Flag Panel to detect irregular characters.
2. 47 billion current and archived web pages.

A handwritten signature in black ink, consisting of a stylized 'S' followed by a cross-like mark.



Turnitin, LLC  
2101 Webster St Ste 1800  
Oakland CA 94612 USA

**RE: Proprietary Certificate**

To Whom It May Concern:

This letter confirms that Turnitin, LLC, having its principal office at Oakland, California, USA, is the, "Sole Source" for the cloud-based originality-checking software service iThenticate. The Turnitin websites, [www.turnitin.com](http://www.turnitin.com), [www.ithenticate.com](http://www.ithenticate.com), and services are proprietary products of Turnitin LLC.

Yours Sincerely,

A handwritten signature in black ink, appearing to be 'Ar' or 'Angela'.

Angela Rhee for Chris Caren  
Chief Executive Officer, Turnitin LLC



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Turnitin, LLC  
2101 Webster St Ste 1800  
Oakland CA 94612 USA

Dear Sirs:

This letter is to attest that Turnitin India Private Limited, a Turnitin group company, exclusively distributes Turnitin LLC's software services – Feedback Studio, Originality Check, iThenticate, SimCheck, and Turnitin Similarity – in India, Nepal, Bangladesh, Bhutan, Sri Lanka, and The Maldives. Please feel free to contact me for further information.

Yours Sincerely,

A handwritten signature in black ink, appearing to be 'Ar' or similar, written over a horizontal line.

Angela Rhee for Chris Caren  
Chief Executive Officer, Turnitin LLC



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**Kalyan Singh Super Specialty Cancer Institute**

कल्याण सिंह अति विशिष्ट कैंसर संस्थान  
C.G. City, Sultanpur Road, Lucknow-226002  
सी०जी० सिटी, सुल्तानपुर रोड, लखनऊ- 226002  
(An Autonomous Institute of the Govt. of Uttar Pradesh)  
(उत्तर प्रदेश सरकार का स्वायत्तशासी संस्थान)

Email: jdmm.sscih@gmail.com

**Proprietary Article Certificate (PAC) for Items/Goods**

(1) The indented goods are manufactured by  
M/s. HemoCue AB

(2) No other make or model is acceptable for the following reasons:

- a) No other hemoglobin analyser can measure
- b) Hb in the range of 0.3 to 30 g/L
- c) No other hemoglobin analyser can measure  
Hb in supernatant plasma.

TZ  
Dr. Brijesh Kumar Yadav  
(Signature of Indentor)

Antu  
24.6.24  
DR. ANTU DUBEY  
(Signature of HOD)





8

Ängelholm  
January 31, 2019

#### Proprietary Certification

Whereas, HemoCue AB, who are established and reputable manufacturers of

HemoCue® Hb 201\* Analyzer  
HemoCue® Hb 201 Microcuvette  
HemoCue® Hb 301 Analyzer  
HemoCue® Hb 301 Microcuvette  
HemoCue® Glucose 201\* Analyzer  
HemoCue® Glucose 201 Microcuvette  
HemoCue® Glucose 201 RT Analyzer  
HemoCue® Glucose 201 RT Microcuvette  
HemoCue® Plasma/Low Hb Photometer  
HemoCue® Plasma/Low Microcuvette  
HemoCue® Albumin 201 Analyzer  
HemoCue® Urine Albumin Microcuvette  
HemoCue® WBC Analyzer  
HemoCue® WBC Microcuvette  
HemoCue® WBC DIFF Analyzer  
HemoCue® WBC DIFF Microcuvette

having factories at

Kuvettgatan 1  
SE-262 71 Ängelholm,  
Sweden

do hereby certify that HemoCue® Plasma/Low Hb Photometer is proprietary product of HemoCue AB.

We market these products through our authorized channel partner in the Indian Market.

HemoCue AB

Pia Werndrup, Ph.D  
IPR Counsellor

Box 1204  
SE-262 23 ÄNGELHOLM  
SWEDEN

HemoCue AB  
Visit: Kuvettgatan 1  
SE-262 71 Ängelholm  
Sweden  
hemocue.com

Post: HemoCue AB  
Box 1204  
SE-262 23 Ängelholm  
Sweden



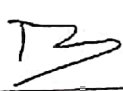
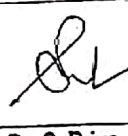
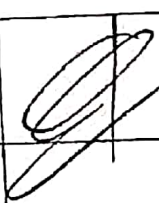
Phone: +46 431 48 12 00  
Fax: +46 431 48 12 25  
E-mail: info@hemocue.se  
Org no: 556342-9272



Eqp	Plasma/Low Hb Analyzer
Qty	01
1.	System should consist of a factory calibrated analyzer and a disposable microcuvettes
2.	The microcuvettes should be made of polystyrene plastic containing sodium deoxycholate, sodium azide, sodium nitrite and nonreactive ingredients
3.	Calibration: Factory calibrated against ICSH reference method. Needs no further calibration
4.	Measuring Range : 0 to 3.0 g/dL
5.	Measurement at 570 & 880nm to compensate for turbidity
6.	Analyzer should carry CE mark.
7.	Analyzer should be US FDA certified
8.	Manufacturer should have ISO 9001:2008 certification
9.	Analyzer Complies with IVD Medical Device Directive 98/79/EC
10.	Measuring Time : One minute
11.	Sample Material : Plasma, serum, aqueous solutions or stored erythrocyte suspensions
12.	Sample Volume : 20 $\mu$ L

One year warranty -

*Any*

				
Dr. Devashish Shukla	Dr. Anju Dubey	Dr. Brijesh Kumar Yadav	Dr. S. Priya	
Medical Superintendent	Associate Prof. Transfusion Med	Assistant. Prof. Transfusion Med	Nominee of Director	F.O./Nominee