BIL	Quotation Reference	Description	Advertisement Date	Closing Date (Not Later Than 09:00AM)	Quotation Fee	Requesting Department	Focal Person
3	PPS/QTN/53/2025)	1/ DISOPYRAMIDE 250MG MODIFIED RELEASE ORAL SOLID PREPARATION PREFERABLY SCORED TABLET. 2/ HYDROCORTISONE 0.5% OINTMENT.	06/05/2025	24/05/2025	\$5.00	JABATAN PERKHIDMATAN FARMASI KEMENTERIAN KESIHATAN	LENNY MARLIANI BINTI HAJI RAMLI  PHARMACIST DRUG  PURCHASING SECTION  TEL: 2393298 ext. 228

QTN REF: PPS/QTN/ 53 /2025

#### SUPPLY AND DELIVERY OF MEDICINES FOR THE DEPARTMENT OF PHARMACEUTICAL SERVICES

NO	DESCRIPTION OF ITEM	QUANTITY	BRAND	MANUFACTURER AND COUNTRY OF ORIGIN	PRICE AND PACK SIZE	TOTAL PRICE
1.	Disopyramide 250mg modified release oral solid preparation preferably scored tablet	20 x 60's				
2.	Hydrocortisone 0.5% ointment	1,800 x 15g - 30g				
First ord upon rec	Y PERIOD: er 2 months, subsequent order 1 month ceipt of purchase order ND LIQUIDATED DAMAGES:					

If the Supplier fails or is unable to deliver the Goods or any parts thereof on the Delivery Date within the time specified, the Government shall be entitled, without prejudice to claim from the Supplier by way of liquidated damages for each day of such delay, a sum of equal to one percent (1%) of the price of the Goods as stated in the relevant Purchase Order in respect of the delayed delivery, provided that the total liquidated damages shall not exceed ten percent (10%) the Purchase Order Price.

#### PRICE VALIDITY:

The quotation shall remain valid for 6 MONTHS from the final date for the submission of the quotation, during which no supplier may withdraw his/her quotation. Where the price validity differs from that required by the Government i.e. 6 months, the **LONGER VALIDITY PERIOD** will be taken as the final validity period.

QTN REF: PPS/QTN/ 53 /2025

#### SUPPLY AND DELIVERY OF MEDICINES FOR THE DEPARTMENT OF PHARMACEUTICAL SERVICES

	TERMS AND CONDITIONS				
a.	Vendor must be registered with the Ministry of Health	Acknowledgement:	Company's Official Stamp		
b.	Please complete the QUOTATION FORM including the USER REQUIREMENT FORM. Submission of incomplete forms may cause DISQUALIFICATION OF QUOTATION	Company Ref. No.:  I hereby certify the above quote to be			
С.	Each vendor is required to quote ONE BRAND WITH ONE PRICE ONLY for each item.	correct. Signature:			
d.	Delivery Period:  FIRST ORDER 2 MONTHS, SUBSEQUENT ORDER 1 MONTH UPON RECEIPT OF PURCHASE ORDER	Name:			
e.	PRICE VALIDITY: The quotation shall remain valid for 6 MONTHS from the final date for the submission of the quotation, during which no supplier may withdraw his/her quotation. Where the price validity differs from that required by the Government i.e. 6 months, the LONGER VALIDITY PERIOD will be taken as the final validity period.	Designation:  Date:			
f.	LETTER OF UNDERTAKING (LOU):  If any of the Goods to be supplied have an expiry date of less than 18 months upon delivery, vendor is required to provide letter of undertaking. The Supplier hereby				

	undertakes to: (i) replace any of the Goods with fresh, new	
No. of the same of	stock; or (ii) issue credit note equivalent to the value of the	
NOT-MATERIAL PROPERTY.	expired Goods.	
g.	Please do not use correction tape or pen/fluid for amendment	

## SAMPLE SUBMISSION FORM

Date:

To:

The Pharmacist

Pharmacy Procurement Section, 3rd Floor, Department of Pharmaceutical Services Building Spg 433, Rimba Highway, Kg Madaras Your Ref: Ministry of Health, Negara Brunei Darussalam				
SUBN	MISSION OF SAMPLES FOR QUOTATION REF	: PPS / QTN / 53	/ 2025	
	nit sample to Pharmacy Procurement Section of quotation advertisement.	no later than FOL	IR WEEKS after g	closing
	ring to the above quotation reference, please sec table below.	e our feedback on	the sample subm	nission
NO	ITEM	SAMPLE SUBMITTED (indicate with v)	SAMPLE NOT SUBMITTED (indicate X with reason)	
1/	Disopyramide 250mg modified release oral solid preparation preferably scored tablet	umania di manine i matemia este del tribune del di matempleta (mente del matempleta) del matempleta (mente d	And the state of t	
2/	Hydrocortisone 0.5% ointment		***************************************	
Thank	you.	Compan	y's Official Stamp	
Name Position Comp	on:			
	FOR OFFICIAL	USE		
Samp	le received by:			
Date :	eceived:			

QTN REF: PPS/QTN/53/2025

No	Requirements	Enter Response Here
1	Sample	Enter Response Here
,	Vendor is required to submit sample in untampered original pack including package insert and Summary of Product Characteristics (in English)  For Controlled drugs and Psychotropics: In the event that sample cannot be submitted, photos and/or artwork are required. See 'Presentation'.	
2	Presentation	
	Vendor is to submit:  i. Clear colour-printed photo of the product offered with company's official stamp. Photo must show label details of the primary and secondary packaging from all sides/angles including name / brand of item, strength and form / preparation.	
	ii. High resolution photo of the following  • For tablets / capsules:  • Appearance of individual tablets  / capsules;  • If the item is in strip pack, the back and front of the strip  • For Injections:  • Appearance of individual vial / ampoule / syringe	
3	Registration with Brunei Darussalam Medicines Control Authority (BDMCA)	
	A copy of any of the following:	
	Priority will be given to medicinal products already registered with the BDMCA.	

4	Manufacturer details	
	Please provide manufacturer details with supporting documents.	
	If manufacturer details are not available, please provide the following:	
	<ul> <li>i. Hardcopy of declaration letter from the principal to inform any issues related to product safety, quality and efficacy as a result of a recall by the manufacturer, wholesaler or Department of Health</li> <li>ii. A copy of the principal's wholesaler license.</li> </ul>	
5		
	Shelf life	
	Please indicate the product shelf-life. Priority will be given to product with a minimum of 24 months.	
6	Storage condition	
	The storage labelling should be in accordance with the latest guideline on registration of medicinal products in Brunei Darussalam.	
	Priority is given to products with specified storage conditions. Terms such as "ambient conditions", "room temperature" or "does not require any special storage condition" should be avoided unless stability studies are provided.	
7	Alcohol and animal content	
	Declaration of source of animal origin and/or alcohol content (if any) is to be provided.	
8		
	Certificate of Analysis	
	A copy of the product's Certificate of Analysis (CoA) is to be submitted.	
	•	

9	New Product	
	Where the product offered has never been supplied to the Ministry of Health, Brunei, detailed information on the product is to be submitted. The information required include, but not limited to, the following:	
The second secon	<ul> <li>i. Bioequivalence studies (Generic products) and / or Clinical studies</li> <li>ii. Stability studies</li> <li>iii. Certificate of free sales</li> <li>iv. Certificate of Pharmaceutical Product (CPP)</li> <li>v. A copy of the Summary of Product Characteristics</li> <li>vi. Good Manufacturing Practice (GMP) certificate</li> <li>vii. Batch release certificate or certificate of origin (for blood products)</li> </ul>	
10	Price Justification	AND CONTRACTOR OF THE PROPERTY
10		
	Vendor is to submit letter of justification on price increase if the same product has been previous supplied to Ministry of Health from the same supplier / distributor.	
11	Local content & Tax Compliance Certificate	
a income and a second a second and a second	Vendor is to provide a copy of the latest content of the company as well as the updated tax compliance certificate, if applicable	
12	Product Registration Number in any of the *benchmark/reference countries	
	If not registered with BDMCA, priority will be given to products which are registered in any of the benchmark/reference countries.	
Note that the state of the stat	*The benchmark/reference countries are Australia, Canada, Malaysia, Singapore, United Kingdom, the United States of America, France, Sweden, Japan, Switzerland, Republic of Korea & European Union.	

### 13 Patent Declaration

- i. Vendors quoting generic products must provide an official declaration or confirmation from the Brunei's Intellectual Property Office verifying whether the innovator product is off-patent
- ii. Vendors quoting innovator products must provide information on the patent status and expiry in Brunei.