



## **Under 16 Cancer Patient Experience Survey 2020**

## Cambridge University Hospitals NHS Foundation Trust



### Contents

Executive Summary	3
	5
Methodology       Scoring methodology         Scoring methodology       Scoring methodology         Eligibility, fieldwork and survey methods       Suppression	5 6 5 6
Understanding the results	7
Further information	7
About the respondents	8
PTC results	<ol> <li>11</li> <li>13</li> <li>13</li> <li>14</li> <li>16</li> <li>17</li> <li>19</li> <li>20</li> <li>21</li> </ol>
Sub-group comparisons       Gender         Gender       Survey type         Survey type       Deprivation         Diagnostic group       Diagnostic group	22 22 23 23 24 25
Benchmarking Charts	26 31



### **Executive Summary**

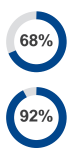
#### **Overall PTC response rate**

81 responded out of a total of 207 eligible parents and children who were sent a survey, resulting in a response rate of 39%. Please note that a response consists of one survey completion, which could consist of both parent and child responses.

	Original sample size	Adjusted sample size <sup>1</sup>	Completed	Response rate
PTC	208	207	81	39%

#### **Overall PTC care rating**

Data for questions in which the base size per question was <11 have been suppressed and replaced with an asterisk (\*).



Children are looked after very well for their cancer or tumour by healthcare staff (Question X65)

Parents or carers rated the overall experience of their child's care as 8 or more out of 10 (Question X64)

<sup>&</sup>lt;sup>1</sup>The adjusted sample excludes patients who were discovered to be ineligible during fieldwork.



#### PTC key question scoring

The following questions were rated as the most important items for providing a good patient experience by professionals involved in children's tumour and cancer care. Scores for all questions can be found in the PTC Excel Data Tables, available on the survey website: <u>https://www.under16cancerexperiencesurvey.co.uk/technical-reports</u>

Data for questions in which the base size per question was <11 have been suppressed and replaced with an asterisk (\*).





#### Introduction

- The Under 16 Cancer Patient Experience Survey 2020 is the first iteration of a national survey that is expected to run on an annual basis to measure children's tumour and cancer care. The survey captures the experiences of children who were aged 8 and above at the start of the fieldwork period, but under 16 at the time of their care, and the parents and carers of children who were aged under 16 at the time of their care.
- The survey has been designed to establish a baseline measure of patient experiences of cancer care
  across England and to enable NHS organisations to assess their performance with other providers. In
  future years, the survey will also allow national trends to be tracked over time, along with allowing NHS
  organisations to compare their performance with earlier waves of data collection.
- The survey is overseen by an Under 16's National Cancer Patient Experience Survey Advisory Group
  made up of professionals involved in the provision of children's cancer care, charity representatives,
  cancer patients and parents of children with cancer. This group advises on questionnaire development,
  methodology and reporting outputs. The survey is managed by NHS England and NHS Improvement,
  who commission Picker to oversee survey development, technical design, implementation and analysis
  of the survey.

### Methodology

#### Eligibility, fieldwork and survey methods

The sample for the survey included all patients with a confirmed tumour or cancer diagnosis who received inpatient or day case care from NHS Principal Treatment Centres (PTCs) between January 1, 2020 and December 31, 2020, and were aged under 16 at the time of their discharge. It is important to note that the sampling period took place during an unprecedented year for the health and social care sector, with the outbreak of the global Covid19 pandemic. This had an impact on the NHS and may have influenced people's care perceptions and experiences, and therefore how they answered the survey. In addition to considering the pandemic, Principal Treatment Centres should also apply any local knowledge to the interpretation of their findings.

The fieldwork for the survey was undertaken between April and June 2021. One of three versions of the survey were distributed:

- The 0-7 questionnaire; sent to parents/carers of patients aged between 0 and 7 years old immediately prior to survey fieldwork (30th March 2021)
- The 8-11 questionnaire, sent to parents/carers of patients aged between 8 and 11 years old old immediately prior to survey fieldwork (30th March 2021)
- The 12-15 questionnaire; sent to parents/carers of patients aged between 12 and 15 years old old immediately prior to survey fieldwork (30th March 2021)

Survey version was assigned based on the patient's age immediately prior to survey fieldwork as opposed to their age at the time they received care, to ensure the most age-appropriate version was sent. For instance, there were small differences in survey design, wording and the way that answer options were presented in the 8-11 and 12-15 questionnaire versions.

Questionnaires sent to those aged 8-11 and 12-15 contained a section for the child to complete, followed by a separate section for their parent or carer to complete. Where a child was aged 0-7, the questionnaire was completed entirely by their parent or carer.

The survey asked recipients to answer about their (or their child's) cancer care over the last year. Some patients may have been 16 or 17 years old at the time they received the questionnaire if they were 15 years old at the time of their discharge but then had a birthday or two prior to the survey being sent out.

The survey used a mixed mode methodology. Questionnaires were sent by post and addressed to the parent or carer of the child, with two reminders sent to non-responders, and also included an option to complete the questionnaire online. A Freephone helpline and email address were available for respondents to opt-out, ask questions about the survey, enable respondents to complete their questionnaire over the



phone and provide access to a translation and interpreting facility for those whose first language was not English.

#### **Suppression**

Suppression is used to protect the confidentiality of respondents in instances in which the base size (number of respondents answering a question) is low and there is a chance that the respondent could be identified.

#### **Question-level suppression**

Data for questions in which the base size per question was <11 have been suppressed and replaced with an asterisk (\*).

#### **Double suppression**

Results for any sub-group breakdown adhere to the same suppression level as the question-level suppression but have an additional suppression rule. Where any of the groups within the sub-group breakdown had <11 respondents then the figure for this particular group was suppressed and replaced with an asterisk (\*). If there was only one group within the sub-group that had <11 respondents and was therefore suppressed, the group with the next lowest number of respondents (regardless of whether it was greater than or less than 11) was suppressed and replaced with an asterisk (\*). This rules applies to scores and proportions.

#### Scoring methodology

For the benchmarking charts and score tables sections, a score has been created for questions that address performance in relation to patient experience (this applies to most survey questions, excluding filter questions and demographic questions such as gender or ethnicity). The score shows the percentage of respondents who gave the most favourable response to a question. Any response options that are not applicable are removed before the score is calculated. This summary score can be used to monitor results over time, and to show how organisations compare to one another, and/or to the national average.

From the example table below, the question would be scored as follows:

#### Staff definitely offered parents enough time to make decisions about their child's treatment: 60%

Question text	Answer options	No. of responses	% responses
	Yes, definitely	120	60%
Did staff offer you enough time to make decisions about your child's treatment?	Yes, to some extent	74	37%
	No, but I would have liked this	6	3%
	No, but this was not needed or possible	n/a	n/a

**Important Note:** Scores have been calculated using unadjusted data. In larger samples, scores are ordinarily adjusted to account for the fact that different demographic groups tend to report their experience of care differently. For example, previous analysis indicates that Black and Asian patients report a less positive experience than White patients on patient experience survey questions. Thereby, PTCs with differing populations could potentially lead to results appearing better or worse than they would if they had a slightly different profile of patients. Scores have not been adjusted on the 2020 survey due to small sample size restrictions. Furthermore, PTCs should be mindful that survey responses may be influenced by the type of care provided by PTCs, for example some provide specialised care and treatment.

## PTCs should therefore be mindful of differences in types of care provision, size and possible demographic makeup when comparing their results against the performance of other PTCs.

Full scores to each scored question can be found in the PTC Excel Data Tables, available here: <u>https://www.under16cancerexperiencesurvey.co.uk/technical-reports</u> Meanwhile, more details on scoring can be found in the Technical Document on the survey website (see the '<u>Further information</u>' section below).



#### **Understanding the results**

The 'PTC results' section of this report presents data from some of the survey questions, and shows the percentage of respondents that selected each response option. There is at least one question from each section of the questionnaire presented in a bar chart. Frequency tables for all survey questions (including non-scored questions) can be found in the PTC Excel Data Tables, available here: <u>https://www.under16cancerexperiencesurvey.co.uk/technical-reports</u>

The sub-group comparisons section of this report shows how different groups of respondents answered the overall care questions in the survey (for example by age/survey type, gender, ethnicity and diagnostic group). The benchmarking charts and score tables sections allow you to compare your scored results against the average of all 13 PTCs.

### The percentages in this report have been rounded to the nearest whole percent. Therefore, in some cases the figures may not add up to 100%.

Question numbers relate to the numbering on the data tables, not the questions used on the surveys themselves.

### **Further information**

This research was carried out in accordance with the internal standard for organisations conducting social research (accreditation to ISO27001:2013; certificate number GB10/80275). The 2020 survey data has been produced and published in line with the Code of Practice for Official Statistics.

For more information on development and methodology, please see the Survey Development Report and Technical Document, available on the survey website. For all other outputs, please visit the results section of the website at <a href="https://www.under16cancerexperiencesurvey.co.uk/technical-reports">https://www.under16cancerexperiencesurvey.co.uk/technical-reports</a>



### About the respondents

#### Table 1. Response rate

Please note that a response means one survey completion, which could be completed by both a parent/ carer and a child.

	Original sample size	Adjusted sample size <sup>2</sup>	Completed	Response rate
PTC	208	207	81	39%

#### Table 2. Responses by survey mode

Response mode	n	%
Paper	57	70%
Online	24	30%
Mixed (combination of paper and online) <sup>3</sup>	0	0%
Phone – English	0	0%
Phone – Translation service	0	0%
Total	81	100%

#### Table 3. Responses by survey type

Survey	n	%
0-7	41	51%
8-11	17	21%
12-15	23	28%
Total	81	100%

#### Table 4. Responses by gender of child (Question X66)

Gender of child	n	%
Male	48	59%
Female	32	40%
Prefer not to say	0	0%
Not given	1	1%
Total	81	100%

<sup>&</sup>lt;sup>2</sup>The adjusted sample excludes patients who were discovered to be ineligible during fieldwork.

<sup>&</sup>lt;sup>3</sup>Indicates cases in which the entire parent/carer section was completed in one mode and the entire child section was completed in another mode.



#### Table 5. Responses by ethnicity of child (Question X69)

Ethnicity of child	n	%
White	72	89%
Mixed, Asian, Black	6	7%
Other and Not Given	3	4%
Total	81	100%

#### Table 6. Responses by current care or treatment stage<sup>4</sup> (Question X01)

Stage of care	n	%
Recently diagnosed	0	0%
Watch and wait	7	9%
Currently receiving treatment	34	43%
Finished treatment in last month	4	5%
In remission / long-term follow-up	36	45%
Receiving palliative or end of life care	0	0%
Recently passed away	0	0%
Other	3	4%
Number of respondents	80	100%

#### Table 7. Responses by diagnostic group<sup>5</sup> (from ICD-10 code in patient sample)

Diagnostic group	n	%
Leukaemias, myeloproliferative diseases, and myelodysplastic diseases	32	40%
Lymphomas and reticuloendothelial neoplasms	11	14%
CNS and miscellaneous intracranial and intraspinal neoplasms	16	20%
Other	22	27%
Total	81	100%

#### Table 8. Responses by long-term, unrelated conditions (Question X70)

Other long-term conditions	n	%
Reported unrelated physical or mental health conditions, disabilities or illnesses that have lasted or are expected to last 12 months or more	20	25%
No long-term, unrelated conditions	60	75%
Total	80	100%

<sup>&</sup>lt;sup>4</sup>Based on a select all that apply question.

<sup>&</sup>lt;sup>5</sup>Details of how diagnostic groups were formed can be found in the technical appendix, available on the survey website: <u>https://www.under16cancerexperiencesurvey.co.uk/technical-reports</u>



## Table 9. Responses by main person who answered questions in the children'ssection (Question X68)

Respondent	n	%
The child / young patient	17	44%
The parent or carer	8	21%
Both the child / young patient and the parent or carer together	14	36%
Total	39	100%

## Table 10. Responses by deprivation<sup>6</sup> (based on IMD from postcode in patientsample)

Deprivation	n	%
1 (most deprived)	11	14%
2	13	16%
3	13	16%
4	22	27%
5 (least deprived)	22	27%
Non-England	0	0%
Total	81	100%

<sup>&</sup>lt;sup>6</sup>Indices of Multiple Deprivation (IMD) classifies geographic areas into five quintiles based on relative disadvantage



#### **PTC results**

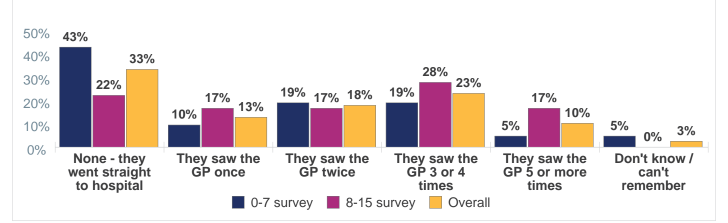
Key findings from each section of the questionnaire can be found below. Please note that full results can be found within the PTC Excel Data Tables (see 'Further information' section for more details).

#### Finding out about the cancer or tumour

49% (n=39) of all parents/carers reported that their children were told they had cancer or tumour within the last year (Question X02). Of this group, 31% (n=12) saw their GP once or twice before being told they needed to go to hospital while 33% (n=13) saw their GP three or more times (Question X04). More details can be found below.

## Figure 1. Answered by parents/carers of all age groups whose children were told they had cancer or a tumour within the last year (Question X04)

Before you were told your child needed to go to hospital about their cancer or tumour, how many times did they see a GP (family doctor) about the health problem caused by the cancer or tumour? (Overall n=39)



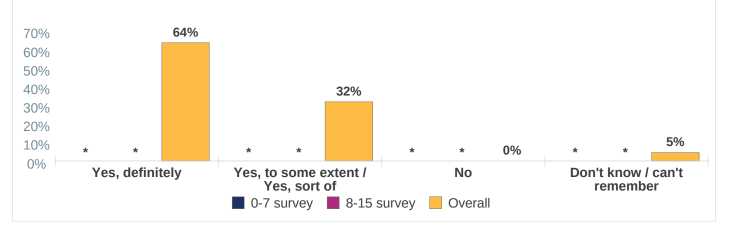
Further questions were asked to all parents/carers of children who had received diagnoses in the past year by the hospital named in the covering letter. Of these parents/carers, 87% (n=20) reported that they were seen as soon as they thought was necessary (Question X07). Meanwhile, 78% (n=18) were definitely told about their child's cancer or tumour in a sensitive way (Question X08) and 65% (n=15) have definitely been able to find the information they needed about their child's diagnosis (Question X12).

Parents/carers of children under the age of 8, along with children aged 8 and above at the time of fieldwork, who were diagnosed in the past year by the hospital named in the covering letter were asked questions concerning how information was given and whether they could have questions answered. Results can be found in the figures below.



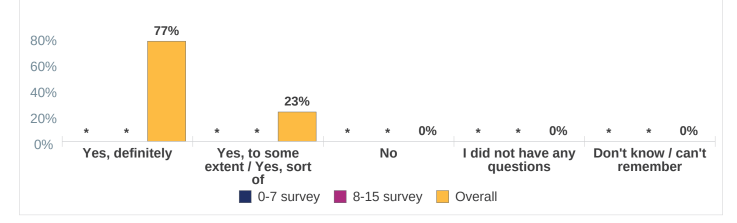
# Figure 2. Answered by parents/carers of 0-7s whose children were told they had cancer or a tumour within the last year & at the hospital named in the letter, and children aged 8-15 who were told they had cancer or a tumour in the last year at the hospital named in the covering letter (Question X09)

When you were told about your child's cancer or tumour, was information given in a way that you could understand? / When you were told about your cancer or tumour, was information given in a way that you could understand? (Overall n=22)



# Figure 3. Answered by parents/carers of 0-7s whose children were told they had cancer or a tumour within the last year & at the hospital named in the letter, and children aged 8-15 who were told they had cancer or a tumour in the last year at the hospital named in the covering letter (Question X10)

Were you able to have any questions answered by healthcare staff after you were told about your child's cancer or tumour? / Were you able to have any questions answered by healthcare staff after you were told about your cancer or tumour? (Overall n=22)

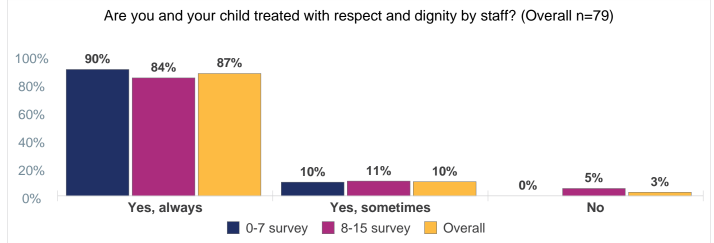




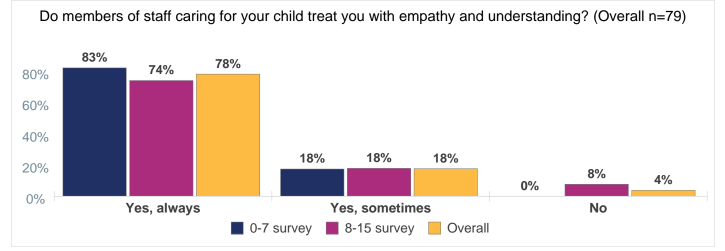
#### Healthcare staff

All parents/carers of children aged under 16 at the time of their care and children aged 8 and above at the time of their care were asked questions about their interactions with healthcare staff at the hospital named in the letter that came with the survey. The results for this section have been broken down into three main themes below: bedside manner and trust, clear communication and support.

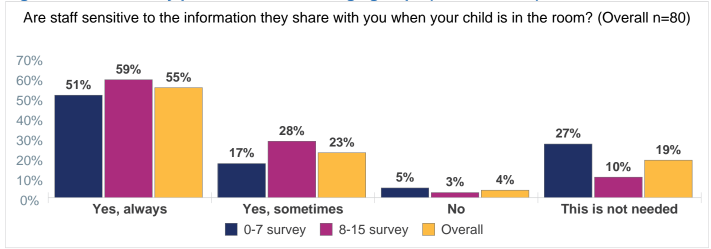
#### **Bedside manner and trust** Figure 4. Answered by parents/carers of all age groups (Question X16)



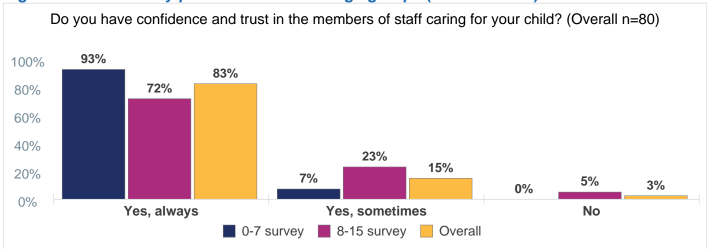
#### Figure 5. Answered by parents/carers of all age groups (Question X18)



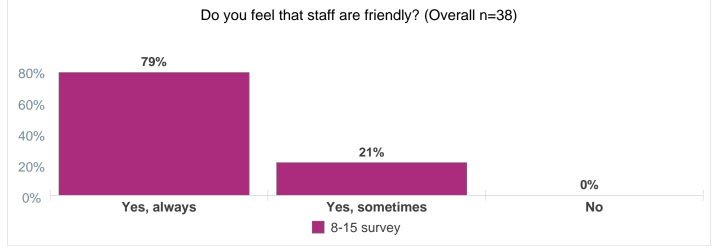
#### Figure 6. Answered by parents/carers of all age groups (Question X20)



#### Figure 7. Answered by parents/carers of all age groups (Question X17)



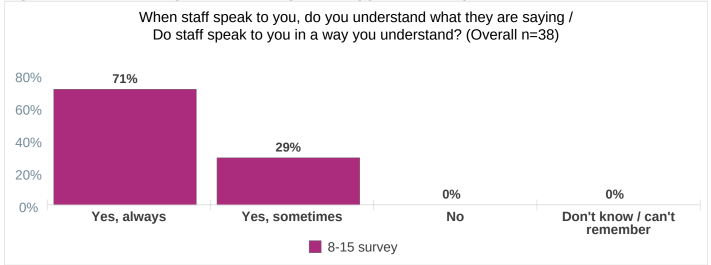
#### Figure 8. Answered by all children aged 8-15 (Question X24)



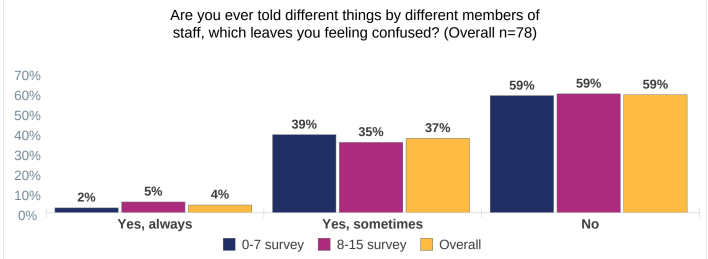
#### **Clear communication** Figure 9. Answered by parents/carers of all age groups (Question X21)

Do healthcare staff share information with your child in a way that is appropriate for them? (Overall n=80) 80% 69% 61% 54% 60% 40% 26% 24% 23% 20% 15% 20% 5% 2% 1% 0% 0% Yes, always Yes, sometimes No This is not needed 0-7 survey Overall

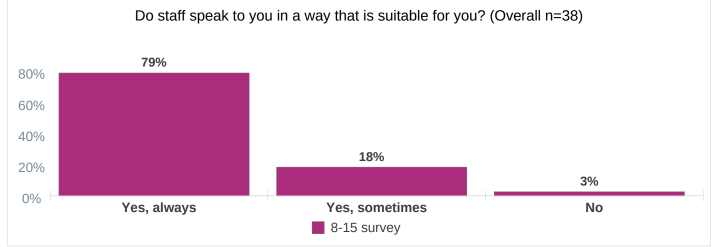
#### Figure 10. Answered by all children aged 8-15 (Question X15)



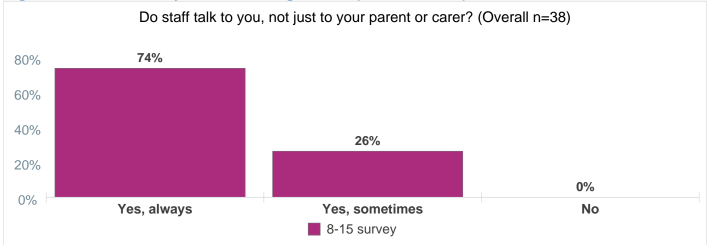
## Figure 11. Answered by parents/carers of children aged 0-7, and all children aged 8-15 (Question X19)



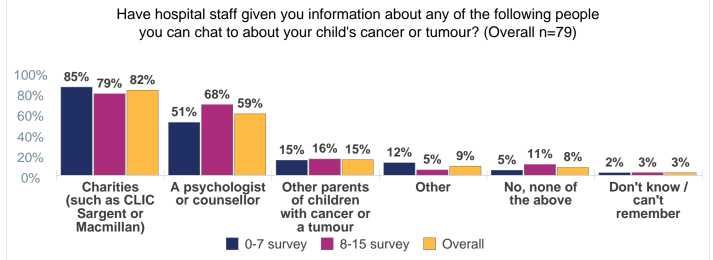
#### Figure 12. Answered by all children aged 8-15 (Question X22)



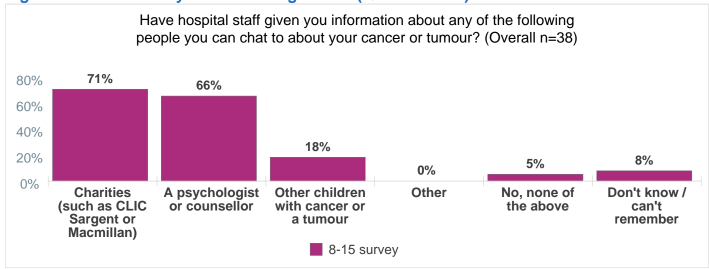
#### Figure 13. Answered by all children aged 8-15 (Question X23)



#### Support Figure 14. Answered by parents/carers of all age groups (Question X26)



#### Figure 15. Answered by all children aged 8-15 (Question X27)

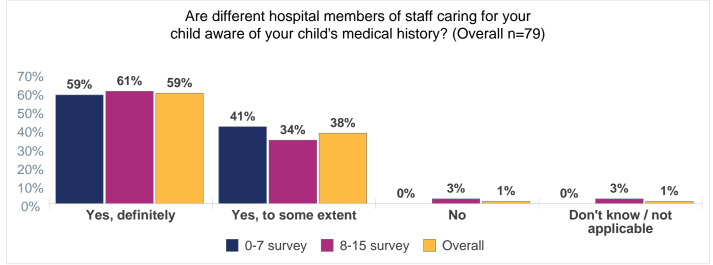




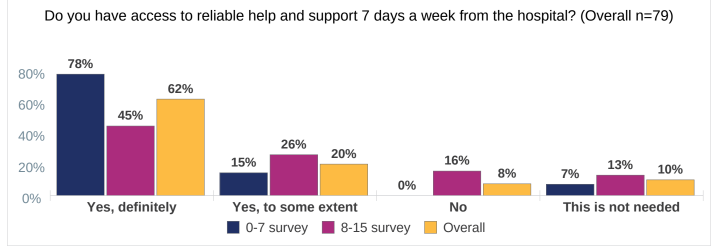
#### **Child's care and treatment**

All parents and carers were asked questions about staff involved in their child's care at the hospital named in the letter that came with their survey, including questions including awareness of the child's medical history and whether they had access to help and support.

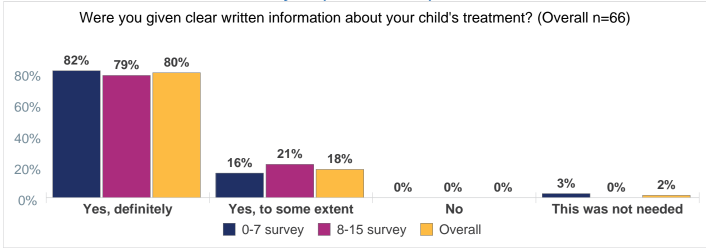
#### Figure 16. Answered by parents/carers of all age groups (Question X30)



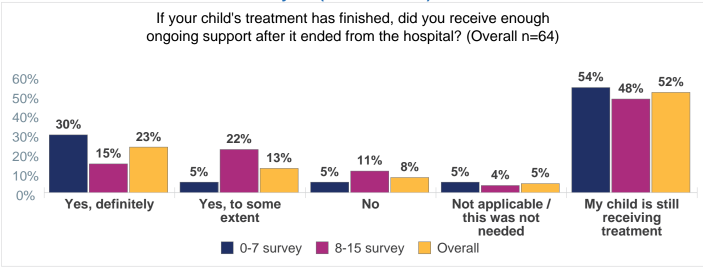
#### Figure 17. Answered by parents/carers of all age groups (Question X35)



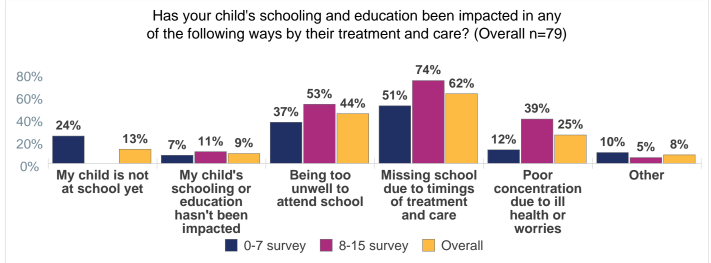
## Figure 18. Answered by parents/carers of all age groups whose children received treatment for their cancer or tumour in the last year (Question X40)



## Figure 19. Answered by parents/carers of all age groups whose children received treatment for their cancer or tumour in the last year (Question X44)



#### Figure 20. Answered by parents/carers of all age groups<sup>5</sup> (Question X33)

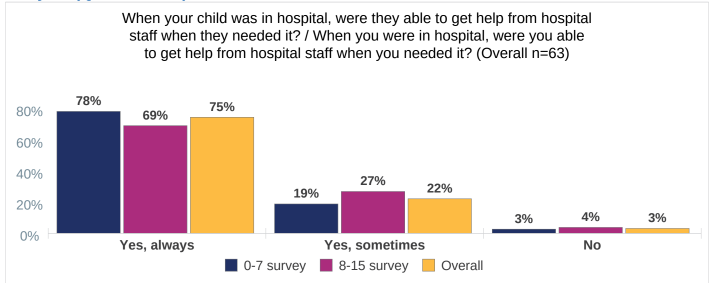




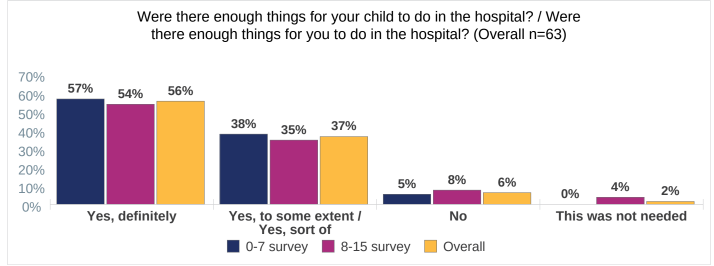
#### Hospital ward

Respondents who had stayed on a hospital ward at the hospital named in the letter that came with their survey (as a day case or for an overnight stay) in the last year were asked questions about hospital staff along with questions regarding services and facilities.

## Figure 21. Answered by parents/carers of children aged 0-7 whose children stayed on a hospital ward in the last year, and children aged 8-15 who stayed on a hospital ward in the last year (Question X47)



## Figure 22. Answered by parents/carers of children aged 0-7 whose children stayed on a hospital ward in the last year, and children aged 8-15 who stayed on a hospital ward in the last year (Question X48)

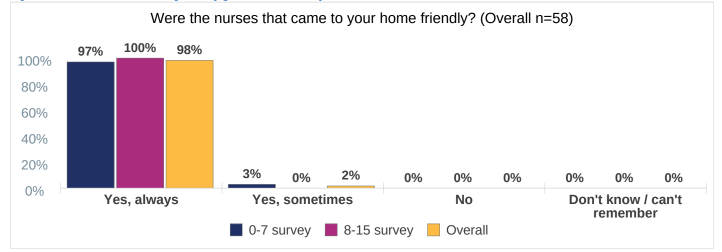




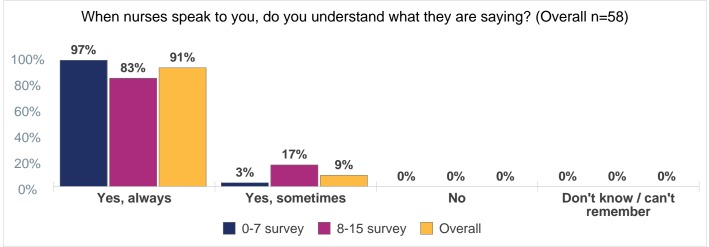
#### Care at home

Children aged 8 and above and parents/carers of children under the age of 8 who had been visited at home by a nurse in the last year, for care relating to the child's cancer or tumour, were asked a short series of questions about care at home. Some results from this section can be found below.

## Figure 23. Answered by parents/carers of children aged 0-7 whose children have been visited at home by a nurse in the last year, and children aged 8-15 who were visited at home by a nurse in the last year (Question X59)



## Figure 24. Answered by parents/carers of children aged 0-7 whose children have been visited at home by a nurse in the last year, and children aged 8-15 who were visited at home by a nurse in the last year (Question X60)



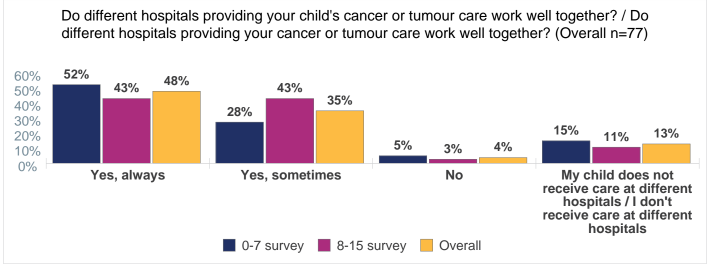


#### **Overall care**

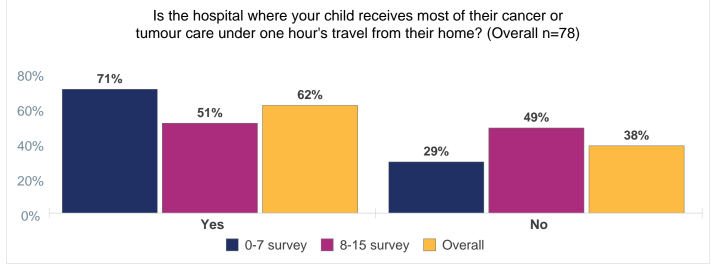
All respondents were asked how they felt about their overall care. Further results for these questions (showing breakdowns by different groups) can be found in the 'Sub-group comparisons' section of this report.

Two additional questions were asked about how well different hospitals providing cancer or tumour care worked together and whether the hospital where the child received most of their cancer or tumour care was under one hours travel time from their home. Results can be found in Figures 25 and 26 below.

## Figure 25. Answered by parents/carers of children aged 0-7 (parent survey), and parents/ carers of children aged 8-11 and children aged 12-15 (children's survey) (Question X62)



#### Figure 26. Answered by parents/carers of all age groups (Question X63)





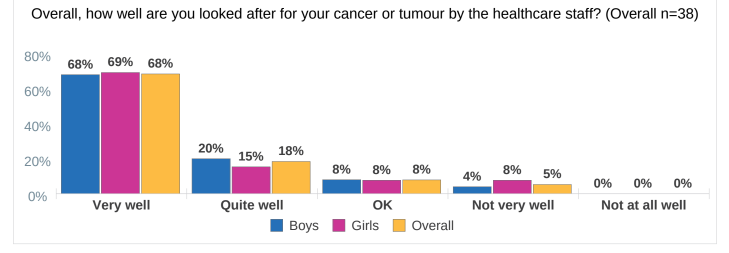
#### Sub-group comparisons

This section summarises the responses of various sub-groups to questions asking about overall care. Further information about how these subgroups were determined can be found in the accompanying technical document, available on the survey website: <u>https://www.under16cancerexperiencesurvey.co.uk/</u>technical-reports.

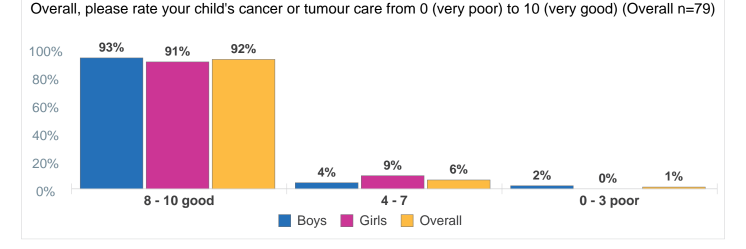
A breakdown of all survey questions by each sub-group can be found in the PTC Excel Data Tables.

Questions asking about overall care were structured differently for children and parents/carers, therefore they cannot be directly compared. Children aged 8 and over were asked how well they were looked after for their cancer or tumour by healthcare staff and were given the options "Very well," "Quite well," "OK," "Not very well" and "Not at all well." Meanwhile, parents and carers of all age groups were asked to rank their child's overall care on a scale of 0-10, with 0 indicating that the care was very poor and 10 indicating that the care was very good. These rankings have been grouped into 0-3 (poor), 4-7, and 8-10 (good) in the results below.

#### **Gender**<sup>7</sup> Figure 27. Answered by all children aged 8-15 (Question X65)



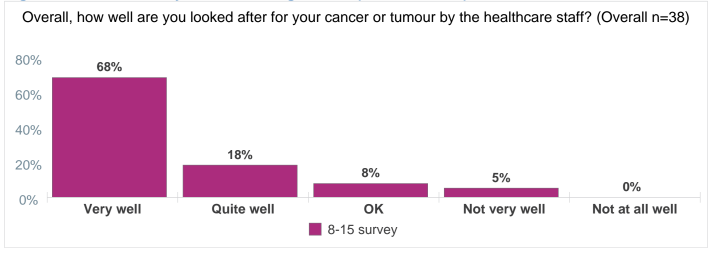
#### Figure 28. Answered by parents/carers of all age groups (Question X64 grouped)



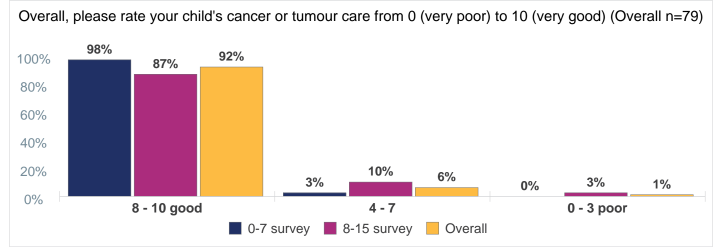
<sup>7</sup>Only data for boys and girls is shown, as the number of respondents answering 'prefer not to say' to the gender question was less than 11.

#### Survey type



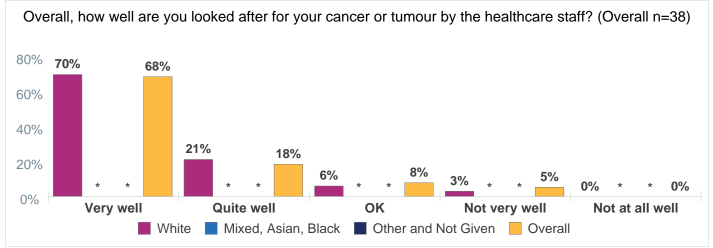


#### Figure 30. Answered by parents/carers of all age groups (Question X64 grouped)

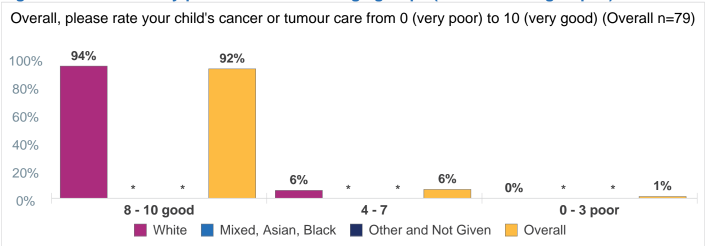


#### Ethnicity

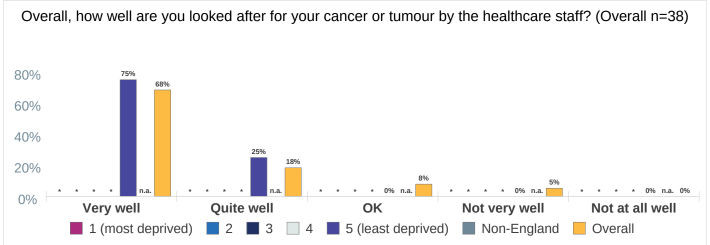
#### Figure 31. Answered by all children aged 8-15 (Question X65)



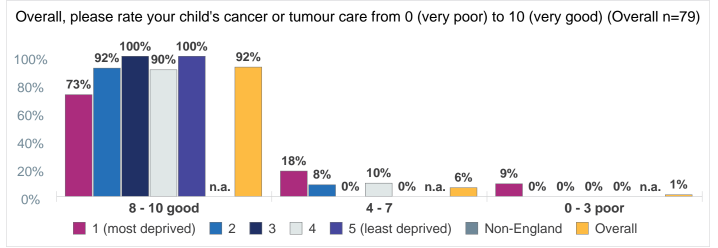
#### Figure 32. Answered by parents/carers of all age groups (Question X64 grouped)



#### **Deprivation** Figure 33. Answered by all children aged 8-15 (Question X65)

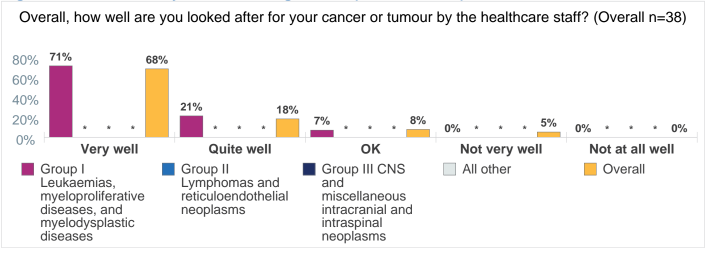


#### Figure 34. Answered by parents/carers of all age groups (Question X64 grouped)

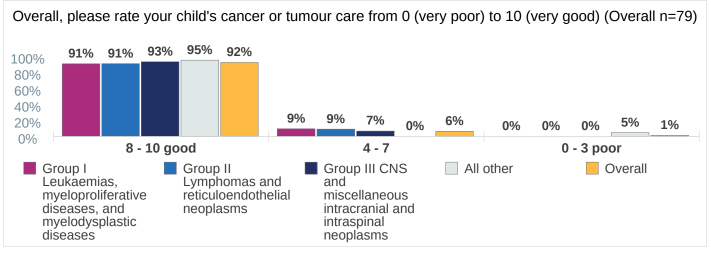


#### **Diagnostic group**





#### Figure 36. Answered by parents/carers of all age groups (Question X64 grouped)





### **Benchmarking Charts**

Benchmarking charts compare experiences in your PTC with those of other PTCs. This allows you to understand where your performance sits in relation to the National average, and compared to the range of scores across all PTCs. Each blue bar shows the range of scores across PTCs, from best performance (to the right) to the poorest performance (to the left). Your PTC score is displayed by the yellow diamond, and the black line shows the National average.

The exact scores for your PTC and for the national average score, for each scored question, can be found in the next section "scoring tables".

**Important Note:** Scores have been calculated using unadjusted data. In larger samples, scores are ordinarily adjusted to account for the fact that different demographic groups tend to report their experience of care differently. For example, previous analysis indicates that Black and Asian patients report a less positive experience than White patients on patient experience survey questions. Thereby, PTCs with differing populations could potentially lead to results appearing better or worse than they would if they had a slightly different profile of patients. Scores have not been adjusted on the 2020 survey due to small sample size restrictions. Furthermore, PTCs should be mindful that survey responses may be influenced by the type of care provided by PTCs, for example some provide specialised care and treatment.

PTCs should therefore be mindful of differences in types of care provision, size and possible demographic makeup when comparing their results against the performance of other PTCs.

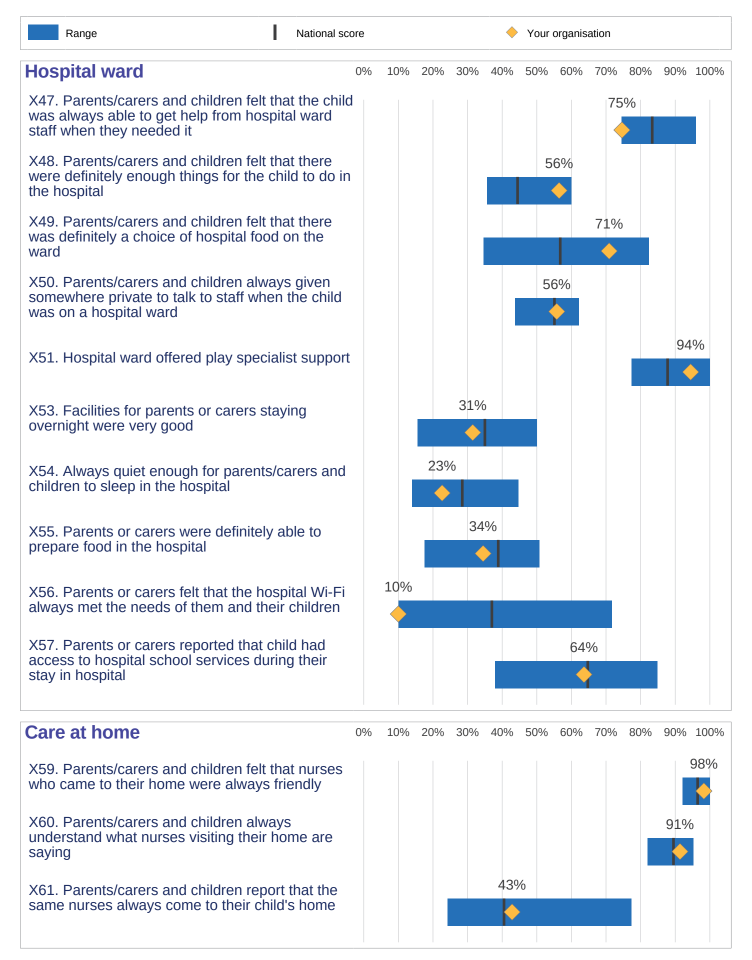
Data for questions in which the base size per question was <11 have been suppressed, and therefore no diamond for your organisation score will be displayed.

Range	National score	Your organisation
Finding out about the cance	er or 0% 10%	20% 30% 40% 50% 60% 70% 80% 90% 100%
tumour		48%
X04. Referral: Saw GP only once or being told they needed to go to hos		87%
X07. Parents or carers thought that was seen as soon as they thought we for their first appointment with a hose	was necessary	78%
X08. Parents or carers definitely tole cancer or tumour in a sensitive way		67%
X09. Information definitely given in a carers and children could understar	a way parents/ nd	77%
X10. Parents/carers and children de have questions answered after bein the cancer or tumour		
X11. Children reported that staff pro about who to contact for more inforr being told about their cancer or turn	mation after	65%
X12. Parents or carers definitely ab information about child's diagnosis	le to find	

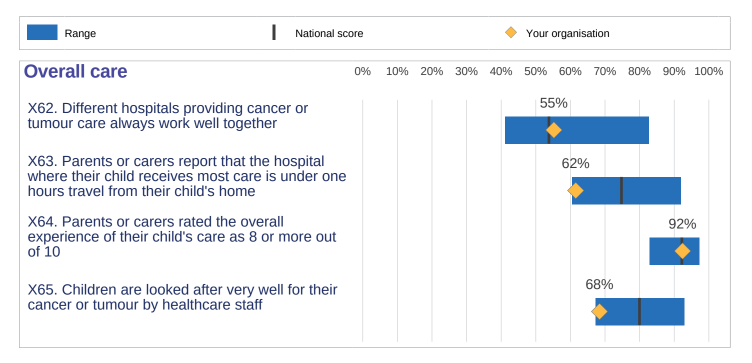
Range	I	National so	core	Your organisation									
Healthcare staff			0%	10%	20%	30%	40%	50%	60%	70%	80%	90%	100%
X13. Staff definitely give parents or information in a way they can under	care stan	rs d									849	%	
X14. Parents or carers definitely had to ask staff questions about child's of treatment											849	%	
X15. Children always understand wi saying	hat s	taff are								71%	,		
X16. Parents or carers feel that they children are always treated with resp dignity by staff											8	87%	
X17. Parents or carers always have and trust in staff caring for their child		fidence									83%	6	
X18. Staff always treat parents or ca empathy and understanding	arers	with									78%		
X19. Parents/carers and children and different things by different members									59%				
X20. Parents or carers feel that staf sensitive to information shared with their child is in the room									6	68% <del>\</del>			
X21. Parents or carers feel that staf share information with children in a appropriate										72%	6		
X22. Child reported that staff always them in a way that is suitable to the		ak to									79%		
X23. Children felt that staff always ta not just their parent or carer	alkec	to them	,							749	%		
X24. Children felt that staff are alwa	ys fri	iendly									79%		
X25. Children reported always or me the same members of staff for their care	ostly treat	seeing ment and	k					53%	6				
X28. Parents or carers have enough about financial help or benefits	n info	ormation									83%	%	

Range	National se	core				•	Your	organis	ation			
Child's care and treatment		0%	10%	20%	30%	40%	50%	60%	70%	80%	90%	100%
X29. Parents or carers feel that differe always work well together	nt staff								69%			
X30. Parents or carers feel that differe definitely aware of their child's medical								60%				
X31. Parents/carers and children alway what is happening with the child's care	ys know								71%			
X32. Parents/carers and children are c involved as much as they want to be in care and treatment		5							69%			
X34. Parents or carers have a named staff at the hospital who can be contact their child's care and treatment											ç	97%
X35. Parents or carers definitely have reliable help and support 7 days a wee hospital									69%			
X36. Parents or carers felt that their ch and treatment is definitely offered at a suitable for their family						40%						
X37. Parents/carers and children felt the child's treatment or care has not change the child is normally able to do, such a friends, doing hobbies or going to school	jed what s seeing				30%							
X38. Parents/carers and children spok about care or treatment changing thing is normally able to do	e with staf is the child	F									94	%
X40. Parents or carers definitely given written information about treatment	clear									82%		
X41. Staff definitely offered parents or enough time to make decisions about treatment	carers child's									79%		
X42. Staff definitely offered support to or carers to help manage child's treatm effects									70%			
X43. Parents or carers definitely able t information they need about their child treatment at the hospital									71%	)		
X44. Parents or carers definitely receiv ongoing support from the hospital after finished							549	%				











#### **Score Tables**

The following tables compare experiences of care in your PTC with the average for all 13 participating PTCs. This allows you to understand where your performance sits in relation to the National average.

**Important Note:** Scores have been calculated using unadjusted data. In larger samples, scores are ordinarily adjusted to account for the fact that different demographic groups tend to report their experience of care differently. For example, previous analysis indicates that Black and Asian patients report a less positive experience than White patients on patient experience survey questions. Thereby, PTCs with differing populations could potentially lead to results appearing better or worse than they would if they had a slightly different profile of patients. Scores have not been adjusted on the 2020 survey due to small sample size restrictions. Furthermore, PTCs should be mindful that survey responses may be influenced by the type of care provided by PTCs, for example some provide specialised care and treatment.

PTCs should therefore be mindful of differences in types of care provision, size and possible demographic makeup when comparing their results against the performance of other PTCs.

Data for questions in which the base size per question was <11 have been suppressed and replaced with an asterisk (\*).

#### Finding out about the cancer or tumour

Question	Scored Text	PTC Score	National Score
X04	Referral: Saw GP only once or twice before being told they needed to go to hospital	48%	57%
X07	Parents or carers thought that their child was seen as soon as they thought was necessary for their first appointment with a hospital doctor	87%	83%
X08	Parents or carers definitely told about cancer or tumour in a sensitive way	78%	79%
X09	Information definitely given in a way parents/carers and children could understand	67%	70%
X10	Parents/carers and children definitely able to have questions answered after being told about the cancer or tumour	77%	82%
X11	Children reported that staff provided details about who to contact for more information after being told about their cancer or tumour	*	91%
X12	Parents or carers definitely able to find information about child's diagnosis	65%	72%

#### Healthcare staff

Question	Scored Text	PTC Score	National Score
X13	Staff definitely give parents or carers information in a way they can understand	84%	87%
X14	Parents or carers definitely had the chance to ask staff questions about child's care and treatment	84%	87%
X15	Children always understand what staff are saying	71%	70%
X16	Parents or carers feel that they and their children are always treated with respect and dignity by staff	87%	89%
X17	Parents or carers always have confidence and trust in staff caring for their child	83%	85%
X18	Staff always treat parents or carers with empathy and understanding	78%	86%
X19	Parents/carers and children are never told different things by different members of staff	59%	64%
X20	Parents or carers feel that staff are always sensitive to information shared with them when their child is in the room	68%	75%
X21	Parents or carers feel that staff always share information with children in a way that is appropriate	72%	79%
X22	Child reported that staff always speak to them in a way that is suitable to them	79%	84%
X23	Children felt that staff always talked to them, not just their parent or carer	74%	79%
X24	Children felt that staff are always friendly	79%	89%
X25	Children reported always or mostly seeing the same members of staff for their treatment and care	53%	61%
X28	Parents or carers have enough information about financial help or benefits	83%	76%



#### Child's care and treatment

Question	Scored Text	PTC Score	National Score
X29	Parents or carers feel that different staff always work well together	69%	70%
X30	Parents or carers feel that different staff are definitely aware of their child's medical history	60%	61%
X31	Parents/carers and children always know what is happening with the child's care	71%	69%
X32	Parents/carers and children are definitely involved as much as they want to be in the child's care and treatment	69%	68%
X34	Parents or carers have a named member of staff at the hospital who can be contacted about their child's care and treatment	97%	92%
X35	Parents or carers definitely have access to reliable help and support 7 days a week from the hospital	69%	66%
X36	Parents or carers felt that their child's care and treatment is definitely offered at a time suitable for their family	40%	45%
X37	Parents/carers and children felt that the child's treatment or care has not changed what the child is normally able to do, such as seeing friends, doing hobbies or going to school	30%	19%
X38	Parents/carers and children spoke with staff about care or treatment changing things the child is normally able to do	94%	95%
X40	Parents or carers definitely given clear written information about treatment	82%	79%
X41	Staff definitely offered parents or carers enough time to make decisions about child's treatment	79%	76%
X42	Staff definitely offered support to parents or carers to help manage child's treatment side effects	70%	75%
X43	Parents or carers definitely able to find information they need about their child's treatment at the hospital	71%	73%
X44	Parents or carers definitely received enough ongoing support from the hospital after treatment finished	54%	57%

#### Hospital ward

Question	Scored Text	PTC Score	National Score
X47	Parents/carers and children felt that the child was always able to get help from hospital ward staff when they needed it	75%	83%
X48	Parents/carers and children felt that there were definitely enough things for the child to do in the hospital	56%	44%
X49	Parents/carers and children felt that there was definitely a choice of hospital food on the ward	71%	57%
X50	Parents/carers and children always given somewhere private to talk to staff when the child was on a hospital ward	56%	55%
X51	Hospital ward offered play specialist support	94%	88%
X53	Facilities for parents or carers staying overnight were very good	31%	35%
X54	Always quiet enough for parents/carers and children to sleep in the hospital	23%	28%
X55	Parents or carers were definitely able to prepare food in the hospital	34%	39%
X56	Parents or carers felt that the hospital Wi-Fi always met the needs of them and their children	10%	37%
X57	Parents or carers reported that child had access to hospital school services during their stay in hospital	64%	65%

#### Care at home

Question	Scored Text	PTC Score	National Score
X59	Parents/carers and children felt that nurses who came to their home were always friendly	98%	96%
X60	Parents/carers and children always understand what nurses visiting their home are saying	91%	89%
X61	Parents/carers and children report that the same nurses always come to their child's home	43%	41%

#### **Overall care**

Question	Scored Text	PTC Score	National Score
X62	Different hospitals providing cancer or tumour care always work well together	55%	54%
X63	Parents or carers report that the hospital where their child receives most care is under one hours travel from their child's home	62%	75%
X64	Parents or carers rated the overall experience of their child's care as 8 or more out of 10	92%	92%
X65	Children are looked after very well for their cancer or tumour by healthcare staff	68%	80%