

Cancer Patient Experience Survey

2023 Results

University Hospitals Plymouth NHS Trust

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Executive summary

Questions above expected range

	Case	mix adjusted s	cores	
	2023 score	Lower expected range	Upper expected range	National score
Q16. Patient was told they could go back later for more information about their diagnosis	88%	81%	87%	84%
Q27. Staff provided the patient with relevant information on available support	95%	89%	93%	91%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	80%	72%	80%	76%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	84%	76%	82%	79%

Questions below expected range

	Case	mix adjusted s	cores	
	2023 score	Lower expected range	Upper expected range	National score
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	82%	85%	92%	88%
Q42_1. Patient completely had enough understandable information about their response to surgery	83%	83%	90%	86%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	78%	80%	90%	85%

Introduction

The National Cancer Patient Experience Survey 2023 is the thirteenth iteration of the survey first undertaken in 2010. It has been designed to monitor progress on cancer care; to provide information to drive local quality improvements; to assist commissioners and providers of cancer care; and to inform the work of the various charities and stakeholder groups supporting cancer patients.

The survey was undertaken by Picker on behalf of NHS England and it was overseen by a national Cancer Patient Experience Advisory Group. This Advisory Group set the principles and objectives of the survey programme and guided questionnaire development. The survey was commissioned and managed by NHS England. The survey provider, Picker, is responsible for designing, running and analysing the survey.

The 2023 survey involved 132 NHS Trusts. Out of 121,121 people, 63,428 people responded to the survey, yielding a response rate of 52%.

Methodology

Eligibility, fieldwork and survey methods

The sample for the survey included all adult (aged 16 and over) NHS patients, with a confirmed primary diagnosis of cancer, discharged from an NHS Trust after an inpatient episode or day case attendance for cancer related treatment in the months of April, May and June 2023. The fieldwork for the survey was undertaken between November 2023 and February 2024.

As in the previous eight years, the survey used a mixed mode methodology. Questionnaires were sent by post, with two reminders where necessary, but also included an option to complete the questionnaire online. A Freephone helpline and email was available for respondents to opt out, ask questions about the survey, enable them to complete their questionnaire over the phone and provide access to a translation and interpreting facility for those whose first language was not English.

Note on question comparability

The questionnaire was redeveloped for the 2021 National Cancer Patient Experience Survey. Year on year comparisons between 2021, 2022 and 2023 are included in this report for most questions. A review of the questionnaire in 2023 saw four changes being made:

- The question text for Q23 and Q42 were amended. These questions are no longer deemed comparable to 2021 and 2022.
- The long-term condition question (Q67) was amended to include "Autism or autism spectrum condition" as a response option. And the "Neurological condition" answer option was updated to include an example condition changing it to "Neurological condition, such as epilepsy". These changes see the answer option "Neurological condition, such as epilepsy" as no longer being deemed comparable to 2021 and 2022.
- The ethnic group question (Q71) was amended to include "Roma" as an answer option. The ethnic group question is still deemed comparable to 2021 and 2022.

Case-mix adjustment

Both unadjusted and adjusted scores are presented in this report. Case-mix adjusted scores allow us to account for the impact that differing patient populations might have on results. By using the case-mix adjusted estimates we can obtain a greater understanding of how a Trust is performing given their patient population. The factors taken into account in this case-mix adjustment are Male/Female/Non-binary/Other, age, ethnicity, deprivation, and cancer type.

Unadjusted data should be used to see the actual responses from patients relating to the Trust. Casemix adjusted data, together with expected ranges, should be used to understand whether the results are significantly higher or lower than national results taking account of the patient mix.

Scoring methodology

Sixty-one questions from the questionnaire are scored as these questions relate directly to patient experience. For all but one question (Q59), the score shows the percentage of respondents who gave

the most favourable response to a question. For Q59, respondents rate their overall care on a scale of 0 to 10, of which the average was calculated for this question's score. The percentages in this report have been rounded to the nearest percentage point. Therefore, in some cases the figures do not appear to add up to 100%.

In 2022, following a review of the scoring methodology, a change was made to the scoring of Q12 such that the response option "No, I was told by letter or email" is no longer considered neutral and is now scored as negative.

Statistical significance

In the reporting of 2023 results, appropriate statistical tests have been undertaken to identify unadjusted scores for which the change over time is 'statistically significant'. A statistically significant difference means that the change in the result is very unlikely to have occurred by chance.

Suppression

Data is suppressed for two reasons: to ensure unreliable results based on very small numbers of respondents are not released, and to prevent individuals being identifiable in the data.

In cases where a result is based on fewer than 10 responses, the result has been suppressed. For example, where fewer than 10 people answered a question from a particular Trust, the results are not shown for that question for that Trust.

For Trusts with an eligible population of 1,000 or fewer, data relating to the respondent and their condition has been suppressed where 5 people or fewer were in a particular category. In instances where only one has been suppressed, the next lowest category has been suppressed to prevent back calculation from the total number of responses.

Additional suppression

Additional suppression happens if only **one** Trust has a score suppressed. If this happens, we will suppress another Trust's results (both the Trust level and sub-group results for the question) based on the next lowest number of respondents for the score. We do this so that the national score cannot be used to work out the score for the individual Trust.

The same rule applies to groups in each sub-group breakdown. For example, if only one Trust has the 85+ age group suppressed for Q25 we will need to suppress another Trust's results for the 85+ age group on Q25. This suppression is based on the 85+ age group with the next lowest number of respondents for Q25.

Understanding the results

This report shows how this Trust scored for each question in the survey compared with national results. It is aimed at helping individual Trusts to understand their performance and identify areas for local improvement. Below is a description of the type of results presented within this report and how to understand them.

Expected range charts

The expected range charts in this report show a bar with the lowest and highest score received for each question nationally. Within this bar, an expected range is given (within the grey bar) and a black diamond represents the actual score for this Trust.

Trusts whose score is above the upper limit of the expected range (in the dark blue) are positive outliers, with a score statistically significantly higher than the national mean. This indicates that the Trust performs better than what Trusts of the same size and demographics are expected to perform. The opposite is true if the score is below the lower limit of the expected range (in the light blue); these are negative outliers. For scores within the expected range (in the grey), the score is what we would expect given the Trust's size and demographics.

Comparability tables

The comparability tables show the 2022 and 2023 unadjusted scores for this Trust for each scored question. The Change 2022-2023 and Change overall columns show whether the scores show a statistically significant variation between years. This is shown between 2022-2023 and as an overall

between 2021-2023. An upwards arrow indicates a statistically significant increase, a downwards arrow indicates a statistically significant decrease and no arrow indicates no statistically significant change.

The adjusted 2023 score will also be presented for each scored question along with the lower and upper expected range and national score. Scores above the upper limit of the expected range will be highlighted dark blue, scores below the lower limit of the expected range will be highlighted light blue, and scores within the lower and upper limit of the expected ranges will be highlighted grey.

Sub-group breakdowns

Unadjusted scores are shown for tumour group, Male/Female/Non-binary/Other, age, IMD quintile, long-term condition status and ethnicity breakdowns. Unadjusted scores for the same sub-group across different Trusts may not be comparable, as they do not account for the impact that differing patient populations might have on results.

Tumour group tables

The tumour group tables show the unadjusted scores for each scored question for each of the 13 tumour groups. Central nervous system is abbreviated as 'CNS' and lower gastrointestinal tract is abbreviated as 'LGT' throughout this report.

Age group tables

The age group tables show the unadjusted scores for each scored question for each of the eight age groups.

Male/Female/Non-binary/Other tables

These tables show the unadjusted scores for the following groups male; female; non-binary; prefer to self-describe; and prefer not to say.

Ethnicity tables

The ethnicity tables show the unadjusted scores for six ethnicity groups.

Long-term condition status tables

The long-term condition status tables show the unadjusted scores for two groups: those who indicate they have one or more long-term conditions and those who indicate that they have no long-term conditions.

IMD quintile tables

The IMD quintile tables show the unadjusted scores for five quintiles based on relative disadvantage, with quintile 1 being the most deprived and quintile 5 being the least deprived.

Year on year charts

The year on year charts show three columns representing the unadjusted scores of the last three years (2021, 2022 and 2023) for each scored question.

National level and England level data

In some cases (343 respondents in 2023), patients from outside England (from Wales, Scotland, Northern Ireland, the Channel Islands or the Isle of Man) are referred to English NHS Trusts for treatment. These patients are described as 'Non-England' in the data.

National level data (England and Non-England) is used for:

- · Response rate section
- National column in comparability tables section
- Sub-group tables section (Tumour group tables, Age group tables, Male/Female/Non-binary/Other tables, Ethnicity tables, IMD quintile tables and Long-term condition status tables)

England only level data is used for:

- Expected range charts section (as case-mix adjustment includes IMD data specific to England)
- Comparability tables section
- Year on year charts section.

Further information

This research was carried out in accordance with the international standard for organisations conducting market and social research (accreditation to ISO20252:2019; certificate number GB08/74322). Our statistical practice is regulated by the Office for Statistics Regulation (OSR). OSR sets the standards of trustworthiness, quality, and value in the Code of Practice for Statistics that all producers of official statistics should adhere to. You are welcome to contact us directly with any comments about how we meet these standards. Alternatively, you can contact OSR by emailing regulation@statistics.gov.uk or via the OSR website.

For more information on the methodology, please see the Technical Document. It can be viewed along with the 2023 questionnaire and survey guidance on the website at www.ncpes.co.uk. For all other outputs at Trust level, please see the Excel tables and dashboards at www.ncpes.co.uk.

Response rate

Overall response rate

883 patients responded out of a total of 1,443 patients, resulting in a response rate of 61%.

	Sample size	Adjusted sample	Completed	Response rate
Overall response rate	1,553	1,443	883	61%
National	129,231	121,121	63,438	52%

Respondents by survey type

	Number of respondents
Paper	706
Online	175
Phone	1
Translation service	1
Total	883

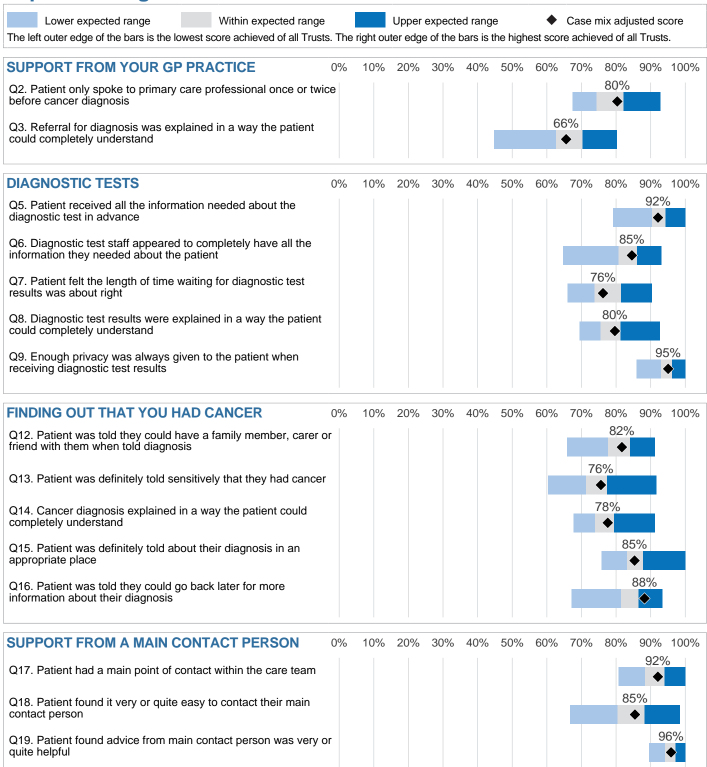
Respondents by tumour group

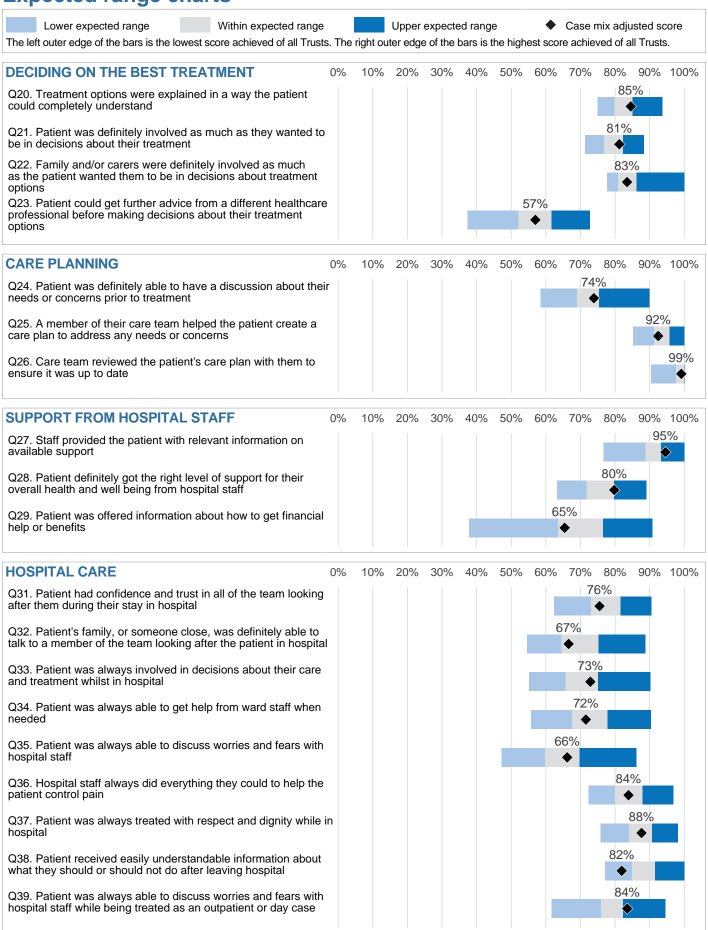
	Number of respondents
Brain / CNS	9
Breast	130
Colorectal / LGT	85
Gynaecological	27
Haematological	120
Head and neck	20
Lung	73
Prostate	106
Sarcoma	8
Skin	64
Upper gastro	59
Urological	71
Other	111
Total	883

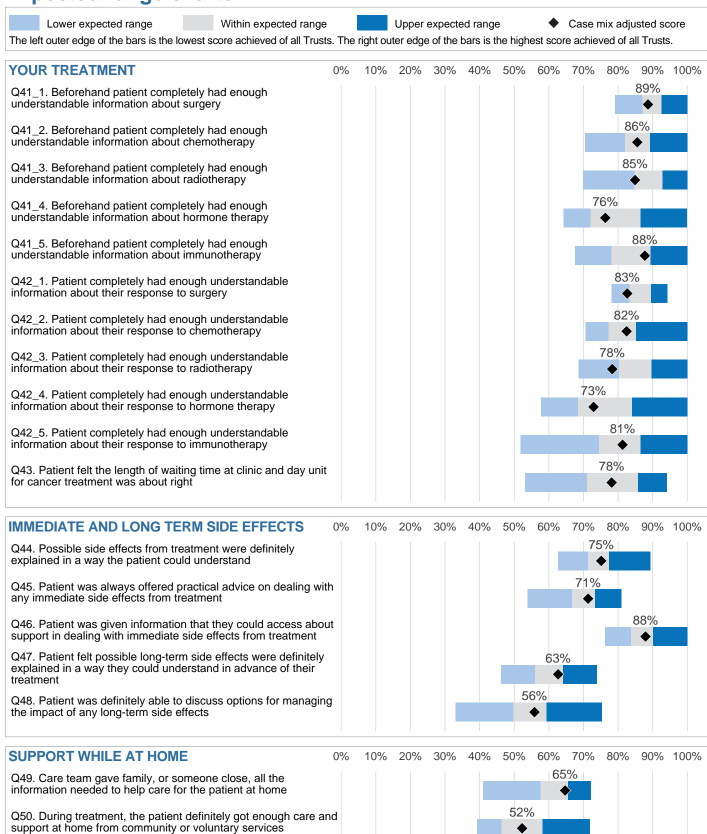
Respondents by ethnicity

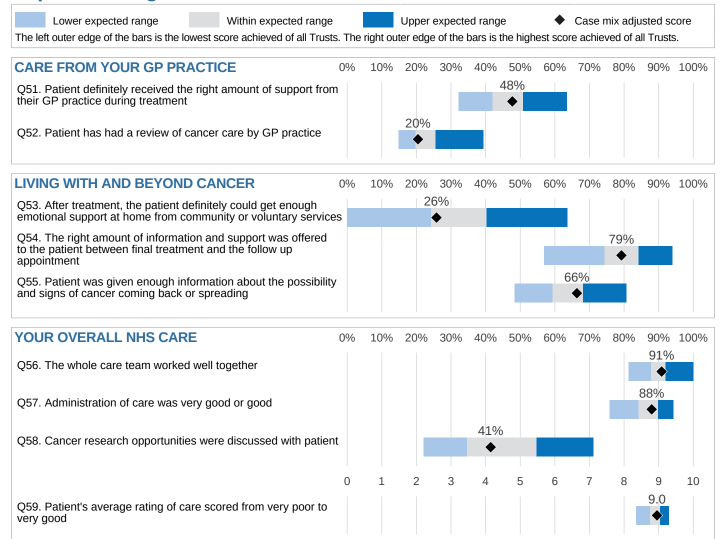
	Number of respondents
White	
English / Welsh / Scottish / Northern Irish / British	801
Irish	*
Gypsy or Irish Traveller	*
Roma	*
Any other White background	11
Mixed / Multiple Ethnic Groups	
White and Black Caribbean	*
White and Black African	*
White and Asian	*
Any other Mixed / multiple ethnic background	*
Asian or Asian British	
Indian	*
Pakistani	*
Bangladeshi	*
Chinese	*
Any other Asian background	*
Black / African / Caribbean / Black British	
African	*
Caribbean	*
Any other Black / African / Caribbean background	*
Other Ethnicity	
Arab	*
Any other ethnic group	*
Not given	
Not given	57
Total	883

^{*} indicates the count is not shown due to suppression









Comparability tables

Indicates where a score is not available due to suppression or a low base size.

** No score available for 2022.

Change 2022-2023: Indicates where 2023 score is significantly higher or lower than 2022 score. Change overall: Indicates significant change overall (2021, 2022, and 2023).

Adjusted Score below Lower Expected Range

Adjusted Score between Upper and Lower Expected Ranges Adjusted Score above Upper Expected Range

	Unadjusted scores Case mix adjusted scores									
SUPPORT FROM YOUR GP PRACTICE	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis		75%	435	79%			80%	74%	82%	78%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	576	67%	578	63%			66%	63%	70%	67%

			Unadjus	ted score	Case n					
DIAGNOSTIC TESTS	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q5. Patient received all the information needed about the diagnostic test in advance	704	92%	713	92%			92%	90%	94%	92%
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	737	85%	757	85%			85%	81%	86%	83%
Q7. Patient felt the length of time waiting for diagnostic test results was about right	737	79%	758	76%		•	76%	74%	81%	78%
Q8. Diagnostic test results were explained in a way the patient could completely understand	737	81%	760	80%			80%	76%	81%	78%
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	739	95%	763	95%			95%	93%	96%	95%

			Unadjus	ted score		Case n				
FINDING OUT THAT YOU HAD CANCER	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	806	75%	805	81%	A	A	82%	78%	84%	81%
Q13. Patient was definitely told sensitively that they had cancer	845	74%	865	75%			76%	71%	77%	74%
Q14. Cancer diagnosis explained in a way the patient could completely understand	849	78%	871	77%			78%	74%	80%	77%
Q15. Patient was definitely told about their diagnosis in an appropriate place	844	84%	864	85%			85%	83%	88%	86%
Q16. Patient was told they could go back later for more information about their diagnosis	755	85%	775	88%			88%	81%	87%	84%

			Unadjust	ed score	Case m					
SUPPORT FROM A MAIN CONTACT PERSON		2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q17. Patient had a main point of contact within the care team	816	93%	846	92%			92%	88%	94%	91%
Q18. Patient found it very or quite easy to contact their main contact person	694	85%	711	86%			85%	80%	88%	84%
Q19. Patient found advice from main contact person was very or quite helpful	717	96%	738	96%			96%	94%	97%	96%

Comparability tables

Adjusted Score below Lower

*	Indicates where a score is not available due to suppression or a low base size. No score available for 2022.	A 0	r ▼	Change 2022-2023: Indicates where 2023 score is significantly higher or lower than 2022 score. Change overall: Indicates significant change overall (2021, 2022, and 2023).	Expected Range Adjusted Score between Upper and Lower Expected Ranges Adjusted Score above Upper Expected Range

			Unadjust	ted score	s		Case n	d scores		
DECIDING ON THE BEST TREATMENT	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q20. Treatment options were explained in a way the patient could completely understand	783	84%	812	84%			85%	80%	85%	82%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	843	81%	853	81%			81%	77%	82%	80%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	699	82%	747	84%		A	83%	81%	86%	83%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	**	**	421	57%			57%	52%	62%	57%

		,	Unadjust	ted score	es		Case n	d scores		
CARE PLANNING	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	756	72%	764	73%			74%	69%	75%	72%
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	433	95%	459	93%			92%	91%	96%	94%
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	350	99%	357	99%			99%	98%	100%	99%

			Unadjust	ted score	es		Case n			
SUPPORT FROM HOSPITAL STAFF	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q27. Staff provided the patient with relevant information on available support	711	92%	747	94%			95%	89%	93%	91%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	843	77%	861	80%			80%	72%	80%	76%
Q29. Patient was offered information about how to get financial help or benefits	352	60%	359	65%			65%	64%	77%	70%

Comparability tables

Indicates where a score is not available due to suppression or a low base size.

** No score available for 2022.

Change 2022-2023: Indicates where 2023 score is

Adjusted Score below Lower Expected Range Jpper

ges per Expected Range

▲ or ▼	Change overall: Indicates significant change overall	and Lower Expected Range
	(2021, 2022, and 2023).	Adjusted Score above Upper

	Unadjusted scores							nix adjuste	d scores	
HOSPITAL CARE	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	347	82%	375	76%			76%	73%	82%	77%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	290	73%	317	67%			67%	65%	75%	70%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	343	66%	374	73%			73%	66%	75%	70%
Q34. Patient was always able to get help from ward staff when needed	337	74%	372	72%			72%	68%	78%	73%
Q35. Patient was always able to discuss worries and fears with hospital staff	327	60%	358	66%			66%	60%	70%	65%
Q36. Hospital staff always did everything they could to help the patient control pain	292	84%	336	84%			84%	80%	88%	84%
Q37. Patient was always treated with respect and dignity while in hospital	344	88%	378	88%			88%	84%	91%	87%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	341	85%	366	82%			82%	85%	92%	88%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	730	82%	731	84%			84%	76%	82%	79%

	Unadjusted scores							nix adjuste	ed scores	
YOUR TREATMENT	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q41_1. Beforehand patient completely had enough understandable information about surgery	473	91%	462	89%			89%	87%	93%	90%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	352	90%	374	86%			86%	82%	89%	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	262	91%	233	85%			85%	85%	93%	89%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	138	86%	119	78%			76%	72%	87%	79%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	139	91%	167	89%			88%	78%	89%	84%
Q42_1. Patient completely had enough understandable information about their response to surgery	**	**	460	83%			83%	83%	90%	86%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	**	**	373	82%			82%	77%	85%	81%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	**	**	226	78%			78%	80%	90%	85%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	**	**	115	75%			73%	68%	84%	76%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	**	**	168	82%			81%	75%	87%	81%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	824	81%	822	78%			78%	71%	86%	78%

Comparability tables

 Indicates where a score is not available due to suppression or a low base size.



Change 2022-2023: Indicates where 2023 score is significantly higher or lower than 2022 score. Change overall: Indicates significant change overall (2021, 2022, and 2023).

Adjusted Score below Lower Expected Range

Adjusted Score between Upper and Lower Expected Ranges Adjusted Score above Upper Expected Range

** No score available for 2022.

			Unadjus	ted score	S		Case n	nix adjuste	d scores	
IMMEDIATE AND LONG TERM SIDE EFFECTS	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	795	77%	802	75%			75%	71%	77%	74%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	753	73%	760	72%			71%	67%	73%	70%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	618	86%	623	88%			88%	84%	90%	87%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	738	61%	750	63%			63%	56%	64%	60%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	622	56%	666	56%			56%	50%	59%	55%

			Unadjus	ted score		Case n				
SUPPORT WHILE AT HOME	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	542	60%	575	65%			65%	58%	66%	62%
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	250	50%	271	52%			52%	46%	58%	52%

		,	Unadjust	ted score		Case n				
CARE FROM YOUR GP PRACTICE	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q51. Patient definitely received the right amount of support from their GP practice during treatment	472	48%	480	47%			48%	42%	51%	46%
Q52. Patient has had a review of cancer care by GP practice	806	18%	819	20%			20%	20%	26%	23%

			Unadjus	ed score	s		Case n	d scores		
LIVING WITH AND BEYOND CANCER	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	154	25%	131	26%			26%	24%	40%	32%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	395	77%	380	79%			79%	74%	84%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	647	66%	664	68%			66%	59%	68%	64%

			Unadjus	ted score	es		Case n	ed scores		
YOUR OVERALL NHS CARE	2022 n	2022 score	2023 n	2023 score	Change 2022- 2023	Change overall	2023 score	Lower expected range	Upper expected range	National score
Q56. The whole care team worked well together	801	92%	811	91%			91%	88%	92%	90%
Q57. Administration of care was very good or good	828	87%	852	88%			88%	84%	90%	87%
Q58. Cancer research opportunities were discussed with patient	517	43%	517	42%			41%	35%	55%	45%
Q59. Patient's average rating of care scored from very poor to very good	814	8.9	843	9.0			9.0	8.8	9.0	8.9

SUPPORT FROM YOUR GP PRACTICE							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	*	96%	80%	80%	54%	85%	77%	91%	*	98%	74%	65%	77%	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	*	85%	62%	56%	51%	67%	51%	76%	*	85%	58%	56%	49%	63%

DIAGNOSTIC TESTS							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q5. Patient received all the information needed about the diagnostic test in advance	*	92%	86%	92%	92%	100%	95%	94%	*	94%	94%	93%	93%	92%
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	*	85%	78%	81%	85%	89%	87%	96%	*	94%	79%	81%	80%	85%
Q7. Patient felt the length of time waiting for diagnostic test results was about right	*	65%	81%	63%	88%	89%	78%	85%	*	65%	77%	76%	73%	76%
Q8. Diagnostic test results were explained in a way the patient could completely understand	*	84%	79%	74%	73%	94%	88%	86%	*	82%	79%	67%	78%	80%
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	*	99%	91%	89%	94%	100%	100%	94%	*	98%	94%	92%	95%	95%

FINDING OUT THAT YOU HAD CANCER							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	*	94%	89%	76%	79%	79%	82%	75%	*	73%	84%	66%	83%	81%
Q13. Patient was definitely told sensitively that they had cancer	*	88%	81%	72%	72%	70%	73%	72%	*	77%	75%	70%	72%	75%
Q14. Cancer diagnosis explained in a way the patient could completely understand	*	86%	79%	73%	66%	90%	82%	80%	*	74%	81%	73%	76%	77%
Q15. Patient was definitely told about their diagnosis in an appropriate place	*	96%	86%	73%	79%	80%	93%	81%	*	92%	85%	79%	79%	85%
Q16. Patient was told they could go back later for more information about their diagnosis	*	89%	88%	76%	82%	90%	90%	96%	*	93%	87%	81%	86%	88%

SUPPORT FROM A MAIN CONTACT PERSO	N						Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q17. Patient had a main point of contact within the care team	*	85%	87%	96%	96%	95%	93%	95%	*	93%	93%	93%	94%	92%
Q18. Patient found it very or quite easy to contact their main contact person	*	91%	83%	91%	84%	94%	84%	77%	*	91%	96%	75%	88%	86%
Q19. Patient found advice from main contact person was very or quite helpful	*	96%	94%	100%	99%	100%	95%	97%	*	96%	98%	90%	95%	96%

DECIDING ON THE BEST TREATMENT							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q20. Treatment options were explained in a way the patient could completely understand	*	85%	80%	85%	84%	95%	92%	85%	*	89%	82%	82%	82%	84%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	*	80%	82%	85%	80%	89%	86%	86%	*	84%	77%	82%	79%	81%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	*	90%	86%	81%	88%	83%	89%	77%	*	78%	78%	73%	85%	84%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	*	50%	57%	43%	49%	*	73%	69%	*	50%	59%	50%	58%	57%

CARE PLANNING							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	*	70%	75%	58%	72%	95%	79%	74%	*	80%	75%	65%	75%	73%
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	*	93%	88%	100%	95%	94%	98%	91%	*	96%	92%	88%	89%	93%
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	*	100%	100%	100%	98%	100%	100%	98%	*	100%	97%	100%	100%	99%

SUPPORT FROM HOSPITAL STAFF							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q27. Staff provided the patient with relevant information on available support	*	97%	92%	92%	95%	100%	93%	97%	*	98%	94%	93%	90%	94%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	*	74%	72%	84%	84%	95%	83%	85%	*	85%	79%	76%	81%	80%
Q29. Patient was offered information about how to get financial help or benefits	*	71%	47%	93%	64%	*	68%	63%	*	50%	81%	63%	59%	65%

HOSPITAL CARE							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	*	79%	69%	69%	87%	91%	84%	79%	*	*	66%	66%	71%	76%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	*	58%	54%	57%	86%	*	72%	77%	*	*	69%	51%	63%	67%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	*	71%	65%	75%	93%	100%	74%	77%	*	*	67%	60%	67%	73%
Q34. Patient was always able to get help from ward staff when needed	*	62%	60%	81%	87%	100%	78%	82%	*	*	57%	68%	67%	72%
Q35. Patient was always able to discuss worries and fears with hospital staff	*	50%	67%	71%	84%	80%	76%	82%	*	*	57%	58%	54%	66%
Q36. Hospital staff always did everything they could to help the patient control pain	*	83%	78%	87%	97%	90%	97%	86%	*	*	76%	76%	81%	84%
Q37. Patient was always treated with respect and dignity while in hospital	*	79%	85%	94%	98%	100%	89%	95%	*	*	87%	80%	77%	88%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	*	79%	77%	67%	91%	90%	91%	85%	*	*	84%	73%	77%	82%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	*	80%	75%	82%	88%	94%	79%	91%	*	91%	76%	86%	81%	84%

YOUR TREATMENT							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	*	89%	86%	86%	91%	92%	91%	95%	*	96%	81%	83%	88%	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	*	78%	90%	88%	92%	*	91%	*	*	*	76%	83%	88%	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	*	80%	83%	*	80%	90%	94%	87%	*	*	82%	94%	87%	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	*	80%	*	*	*	*	*	88%	*	*	*	*	50%	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	*	80%	*	*	86%	*	95%	*	*	89%	*	93%	87%	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	*	83%	80%	79%	83%	92%	79%	95%	*	87%	79%	77%	79%	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	*	75%	79%	88%	84%	*	91%	*	*	*	81%	94%	81%	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	*	79%	67%	*	50%	91%	88%	82%	*	*	87%	84%	70%	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	*	72%	*	*	*	*	*	85%	*	*	*	*	60%	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	*	80%	*	*	81%	*	80%	*	*	89%	*	93%	78%	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	*	67%	84%	77%	64%	94%	82%	91%	*	88%	71%	87%	79%	78%

IMMEDIATE AND LONG TERM SIDE EFFECT	TS						Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	*	70%	71%	85%	75%	95%	75%	81%	*	80%	72%	72%	76%	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	*	66%	66%	84%	76%	100%	71%	70%	*	81%	67%	66%	74%	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	*	88%	90%	92%	85%	100%	92%	86%	*	100%	75%	85%	89%	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	*	61%	60%	50%	48%	79%	66%	77%	*	63%	63%	69%	61%	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	*	55%	46%	36%	54%	88%	64%	63%	*	60%	62%	55%	52%	56%

SUPPORT WHILE AT HOME							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	*	58%	57%	53%	68%	80%	71%	72%	*	69%	70%	71%	58%	65%
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	*	43%	41%	60%	57%	*	70%	75%	*	*	33%	55%	42%	52%

CARE FROM YOUR GP PRACTICE							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q51. Patient definitely received the right amount of support from their GP practice during treatment	*	53%	50%	42%	45%	46%	45%	50%	*	68%	39%	36%	45%	47%
Q52. Patient has had a review of cancer care by GP practice	*	15%	20%	27%	17%	40%	25%	24%	*	20%	21%	21%	17%	20%

LIVING WITH AND BEYOND CANCER							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	*	26%	32%	*	25%	*	31%	29%	*	*	8%	*	29%	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	*	76%	75%	*	78%	100%	84%	81%	*	79%	64%	76%	90%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	*	59%	55%	71%	76%	93%	67%	75%	*	80%	63%	63%	62%	68%

YOUR OVERALL NHS CARE							Tumo	our gro	up					
	Brain / CNS	Breast	Colorectal / LGT	Gynaecological	Haematological	Head and neck	Lung	Prostate	Sarcoma	Skin	Upper gastro	Urological	Other	All
Q56. The whole care team worked well together	*	90%	91%	96%	93%	90%	90%	92%	*	95%	85%	87%	89%	91%
Q57. Administration of care was very good or good	*	90%	81%	84%	88%	95%	92%	85%	*	94%	87%	82%	90%	88%
Q58. Cancer research opportunities were discussed with patient	*	29%	29%	35%	65%	82%	32%	49%	*	39%	31%	32%	44%	42%
Q59. Patient's average rating of care scored from very poor to very good	*	8.9	8.5	8.8	9.2	9.5	9.0	9.0	*	9.2	8.7	8.9	8.9	9.0

SUPPORT FROM YOUR GP PRACTICE					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	*	*	83%	74%	74%	80%	81%	91%	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	*	*	71%	48%	66%	68%	58%	67%	63%

DIAGNOSTIC TESTS			Age						
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q5. Patient received all the information needed about the diagnostic test in advance	*	*	71%	93%	91%	90%	96%	96%	92%
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	*	*	87%	85%	85%	81%	86%	89%	85%
Q7. Patient felt the length of time waiting for diagnostic test results was about right	*	*	44%	67%	74%	76%	82%	85%	76%
Q8. Diagnostic test results were explained in a way the patient could completely understand	*	*	75%	75%	83%	79%	81%	81%	80%
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	*	*	94%	92%	96%	94%	96%	96%	95%

FINDING OUT THAT YOU HAD CANCER					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	*	*	76%	81%	78%	79%	85%	87%	81%
Q13. Patient was definitely told sensitively that they had cancer	*	*	78%	61%	74%	72%	81%	91%	75%
Q14. Cancer diagnosis explained in a way the patient could completely understand	*	*	71%	67%	81%	75%	80%	85%	77%
Q15. Patient was definitely told about their diagnosis in an appropriate place	*	*	78%	83%	84%	82%	90%	85%	85%
Q16. Patient was told they could go back later for more information about their diagnosis	*	*	88%	95%	89%	85%	89%	79%	88%

SUPPORT FROM A MAIN CONTACT PERSO	N				Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q17. Patient had a main point of contact within the care team	*	*	89%	90%	93%	93%	92%	88%	92%
Q18. Patient found it very or quite easy to contact their main contact person	*	*	71%	82%	88%	82%	89%	86%	86%
Q19. Patient found advice from main contact person was very or quite helpful	*	*	80%	85%	97%	95%	99%	100%	96%

DECIDING ON THE BEST TREATMENT					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q20. Treatment options were explained in a way the patient could completely understand	*	*	75%	76%	87%	82%	88%	86%	84%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	*	*	72%	75%	82%	82%	84%	81%	81%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	*	*	79%	79%	89%	81%	86%	79%	84%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	*	*	42%	45%	60%	59%	57%	53%	57%

CARE PLANNING					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	*	*	53%	74%	79%	70%	77%	68%	73%
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	*	*	*	90%	94%	90%	97%	88%	93%
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	*	*	*	100%	99%	98%	100%	100%	99%

SUPPORT FROM HOSPITAL STAFF								Age					
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All				
Q27. Staff provided the patient with relevant information on available support	*	*	88%	90%	96%	95%	95%	85%	94%				
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	*	*	61%	68%	79%	79%	87%	86%	80%				
Q29. Patient was offered information about how to get financial help or benefits	*	*	69%	69%	67%	64%	63%	50%	65%				

HOSPITAL CARE					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	*	*	*	60%	74%	75%	81%	*	76%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	*	*	*	58%	68%	71%	65%	*	67%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	*	*	*	60%	74%	75%	73%	*	73%
Q34. Patient was always able to get help from ward staff when needed	*	*	*	60%	67%	73%	76%	*	72%
Q35. Patient was always able to discuss worries and fears with hospital staff	*	*	*	50%	64%	73%	67%	*	66%
Q36. Hospital staff always did everything they could to help the patient control pain	*	*	*	65%	81%	88%	86%	*	84%
Q37. Patient was always treated with respect and dignity while in hospital	*	*	*	85%	89%	88%	89%	*	88%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	*	*	*	68%	85%	82%	80%	*	82%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	*	*	75%	79%	82%	83%	88%	93%	84%

YOUR TREATMENT					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	*	*	79%	88%	90%	89%	91%	89%	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	*	*	67%	90%	91%	80%	90%	*	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	*	*	*	89%	89%	82%	83%	*	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	*	*	*	*	85%	68%	83%	*	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	*	*	*	67%	90%	87%	95%	*	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	*	*	71%	76%	86%	84%	82%	89%	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	*	*	83%	75%	89%	77%	85%	*	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	*	*	*	83%	84%	82%	66%	*	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	*	*	*	*	68%	67%	82%	*	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	*	*	*	67%	87%	78%	84%	*	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	*	*	63%	78%	76%	81%	77%	94%	78%

IMMEDIATE AND LONG TERM SIDE EFFEC	ΓS								
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	*	*	71%	78%	77%	74%	75%	77%	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	*	*	60%	69%	76%	71%	71%	73%	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	*	*	87%	88%	91%	86%	89%	76%	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	*	*	53%	72%	65%	62%	64%	45%	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	*	*	38%	59%	60%	57%	55%	52%	56%

SUPPORT WHILE AT HOME					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	*	*	50%	58%	67%	65%	64%	79%	65%
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	*	*	*	73%	47%	50%	58%	58%	52%

CARE FROM YOUR GP PRACTICE					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q51. Patient definitely received the right amount of support from their GP practice during treatment	*	*	*	36%	42%	51%	45%	47%	47%
Q52. Patient has had a review of cancer care by GP practice	*	*	12%	20%	24%	20%	18%	22%	20%

LIVING WITH AND BEYOND CANCER					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	*	*	*	*	33%	15%	26%	*	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	*	*	*	87%	81%	73%	86%	86%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	*	*	20%	66%	73%	67%	69%	69%	68%

YOUR OVERALL NHS CARE					Age				
	16 - 24	25 - 34	35 - 44	45 - 54	55 - 64	65 - 74	75 - 84	85+	All
Q56. The whole care team worked well together	*	*	71%	90%	90%	91%	92%	97%	91%
Q57. Administration of care was very good or good	*	*	67%	90%	89%	86%	90%	91%	88%
Q58. Cancer research opportunities were discussed with patient	*	*	47%	47%	42%	45%	43%	0%	42%
Q59. Patient's average rating of care scored from very poor to very good	*	*	8.3	8.7	9.0	9.0	9.0	9.1	9.0

SUPPORT FROM YOUR GP PRACTICE			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	77%	81%	*	*	*	71%	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	64%	63%	*	*	*	65%	63%

DIAGNOSTIC TESTS			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q5. Patient received all the information needed about the diagnostic test in advance	92%	93%	*	*	*	89%	92%
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	81%	88%	*	*	*	85%	85%
Q7. Patient felt the length of time waiting for diagnostic test results was about right	71%	82%	*	*	*	69%	76%
Q8. Diagnostic test results were explained in a way the patient could completely understand	79%	81%	*	*	*	69%	80%
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	95%	95%	*	*	*	95%	95%

FINDING OUT THAT YOU HAD CANCER			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	84%	78%	*	*	*	83%	81%
Q13. Patient was definitely told sensitively that they had cancer	78%	73%	*	*	*	67%	75%
Q14. Cancer diagnosis explained in a way the patient could completely understand	77%	78%	*	*	*	72%	77%
Q15. Patient was definitely told about their diagnosis in an appropriate place	86%	84%	*	*	*	80%	85%
Q16. Patient was told they could go back later for more information about their diagnosis	88%	88%	*	*	*	80%	88%

SUPPORT FROM A MAIN CONTACT PERSO	N		Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q17. Patient had a main point of contact within the care team	90%	95%	*	*	*	86%	92%
Q18. Patient found it very or quite easy to contact their main contact person	87%	84%	*	*	*	88%	86%
Q19. Patient found advice from main contact person was very or quite helpful	97%	95%	*	*	*	97%	96%

DECIDING ON THE BEST TREATMENT			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q20. Treatment options were explained in a way the patient could completely understand	83%	86%	*	*	*	80%	84%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	80%	82%	*	*	*	79%	81%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	85%	83%	*	*	*	76%	84%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	53%	59%	*	*	*	74%	57%

CARE PLANNING		Male/Female/Non-binary/Other					
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	71%	76%	*	*	*	71%	73%
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	91%	94%	*	*	*	91%	93%
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	99%	99%	*	*	*	100%	99%

SUPPORT FROM HOSPITAL STAFF			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q27. Staff provided the patient with relevant information on available support	93%	95%	*	*	*	97%	94%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	77%	84%	*	*	*	69%	80%
Q29. Patient was offered information about how to get financial help or benefits	68%	64%	*	*	*	45%	65%

HOSPITAL CARE			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	72%	79%	*	*	*	76%	76%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	64%	71%	*	*	*	56%	67%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	72%	74%	*	*	*	68%	73%
Q34. Patient was always able to get help from ward staff when needed	68%	74%	*	*	*	82%	72%
Q35. Patient was always able to discuss worries and fears with hospital staff	61%	71%	*	*	*	70%	66%
Q36. Hospital staff always did everything they could to help the patient control pain	83%	85%	*	*	*	88%	84%
Q37. Patient was always treated with respect and dignity while in hospital	83%	91%	*	*	*	95%	88%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	77%	85%	*	*	*	90%	82%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	81%	86%	*	*	*	92%	84%

YOUR TREATMENT			Male/Fema	ile/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	86%	91%	*	*	*	95%	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	85%	86%	*	*	*	83%	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	82%	88%	*	*	*	*	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	72%	86%	*	*	*	*	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	90%	87%	*	*	*	*	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	80%	86%	*	*	*	83%	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	80%	85%	*	*	*	83%	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	75%	81%	*	*	*	*	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	70%	83%	*	*	*	*	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	83%	80%	*	*	*	*	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	72%	84%	*	*	*	85%	78%

IMMEDIATE AND LONG TERM SIDE EFFECT	ΓS		Male/Female/Non-binary/Other				
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	73%	77%	*	*	*	81%	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	71%	73%	*	*	*	63%	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	89%	86%	*	*	*	90%	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	58%	68%	*	*	*	59%	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	51%	62%	*	*	*	49%	56%

SUPPORT WHILE AT HOME			Male/Fema	le/Non-bina	ry/Other		
	Female	Not given	All				
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	61%	70%	*	*	*	57%	65%
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	50%	56%	*	*	*	47%	52%

CARE FROM YOUR GP PRACTICE			Male/Fema	le/Non-bina	ry/Other		
	Female Male Non-binary Prefer to self-describe Prefer not to say Not given						All
Q51. Patient definitely received the right amount of support from their GP practice during treatment	44%	50%	*	*	*	45%	47%
Q52. Patient has had a review of cancer care by GP practice	19%	20%	*	*	*	27%	20%

LIVING WITH AND BEYOND CANCER			Male/Fema	le/Non-bina	ry/Other		
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	25%	31%	*	*	*	8%	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	76%	83%	*	*	*	81%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	62%	73%	*	*	*	63%	68%

YOUR OVERALL NHS CARE		Male/Female/Non-binary/Other					
	Female	Male	Non- binary	Prefer to self- describe	Prefer not to say	Not given	All
Q56. The whole care team worked well together	89%	92%	*	*	*	97%	91%
Q57. Administration of care was very good or good	87%	89%	*	*	*	86%	88%
Q58. Cancer research opportunities were discussed with patient	40%	46%	*	*	*	27%	42%
Q59. Patient's average rating of care scored from very poor to very good	8.9	9.1	*	*	*	8.6	9.0

SUPPORT FROM YOUR GP PRACTICE				Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	80%	*	*	*	*	71%	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	63%	*	*	*	*	68%	63%

DIAGNOSTIC TESTS	Ethnicity								
	White	Mixed	Asian	Black	Other	Not given	All		
Q5. Patient received all the information needed about the diagnostic test in advance	92%	*	*	*	*	91%	92%		
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	85%	*	*	*	*	83%	85%		
Q7. Patient felt the length of time waiting for diagnostic test results was about right	76%	*	*	*	*	75%	76%		
Q8. Diagnostic test results were explained in a way the patient could completely understand	80%	*	*	*	*	71%	80%		
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	95%	*	*	*	*	96%	95%		

FINDING OUT THAT YOU HAD CANCER		Ethnicity						
	White	Mixed	Asian	Black	Other	Not given	All	
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	81%	*	*	*	*	80%	81%	
Q13. Patient was definitely told sensitively that they had cancer	76%	*	*	*	*	69%	75%	
Q14. Cancer diagnosis explained in a way the patient could completely understand	78%	*	*	*	*	67%	77%	
Q15. Patient was definitely told about their diagnosis in an appropriate place	85%	*	*	*	*	83%	85%	
Q16. Patient was told they could go back later for more information about their diagnosis	88%	*	*	*	*	81%	88%	

SUPPORT FROM A MAIN CONTACT PERSO	N			Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q17. Patient had a main point of contact within the care team	92%	*	*	*	*	88%	92%
Q18. Patient found it very or quite easy to contact their main contact person	86%	*	*	*	*	83%	86%
Q19. Patient found advice from main contact person was very or quite helpful	96%	*	*	*	*	93%	96%

DECIDING ON THE BEST TREATMENT				Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q20. Treatment options were explained in a way the patient could completely understand	85%	*	*	*	*	80%	84%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	82%	*	*	*	*	76%	81%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	84%	*	*	*	*	80%	84%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	57%	*	*	*	*	59%	57%

CARE PLANNING	Ethnicity						
	White	Mixed	Asian	Black	Other	Not given	All
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	74%	*	*	*	*	67%	73%
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	93%	*	*	*	*	89%	93%
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	99%	*	*	*	*	100%	99%

SUPPORT FROM HOSPITAL STAFF				Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q27. Staff provided the patient with relevant information on available support	94%	*	*	*	*	98%	94%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	80%	*	*	*	*	75%	80%
Q29. Patient was offered information about how to get financial help or benefits	67%	*	*	*	*	40%	65%

HOSPITAL CARE				Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	76%	*	*	*	*	73%	76%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	67%	*	*	*	*	59%	67%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	73%	*	*	*	*	70%	73%
Q34. Patient was always able to get help from ward staff when needed	70%	*	*	*	*	85%	72%
Q35. Patient was always able to discuss worries and fears with hospital staff	66%	*	*	*	*	72%	66%
Q36. Hospital staff always did everything they could to help the patient control pain	84%	*	*	*	*	91%	84%
Q37. Patient was always treated with respect and dignity while in hospital	87%	*	*	*	*	96%	88%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	81%	*	*	*	*	92%	82%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	84%	*	*	*	*	86%	84%

YOUR TREATMENT				Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	89%	*	*	*	*	89%	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	86%	*	*	*	*	90%	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	84%	*	*	*	*	92%	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	78%	*	*	*	*	*	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	89%	*	*	*	*	*	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	83%	*	*	*	*	79%	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	82%	*	*	*	*	89%	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	78%	*	*	*	*	80%	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	75%	*	*	*	*	*	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	81%	*	*	*	*	*	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	78%	*	*	*	*	82%	78%

IMMEDIATE AND LONG TERM SIDE EFFECT	S			Ethnicity			
	White	Mixed	Asian	Black	Other	Not given	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	75%	*	*	*	*	75%	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	72%	*	*	*	*	64%	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	88%	*	*	*	*	92%	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	63%	*	*	*	*	57%	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	56%	*	*	*	*	56%	56%

SUPPORT WHILE AT HOME	Ethnicity						
	White	Mixed	Asian	Black	Other	Not given	All
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	65%	*	*	*	*	54%	65%
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	52%	*	*	*	*	48%	52%

CARE FROM YOUR GP PRACTICE	Ethnicity						
	White	Mixed	Asian	Black	Other	Not given	All
Q51. Patient definitely received the right amount of support from their GP practice during treatment	46%	*	*	*	*	52%	47%
Q52. Patient has had a review of cancer care by GP practice	20%	*	*	*	*	23%	20%

LIVING WITH AND BEYOND CANCER		Ethnicity					
	White	Mixed	Asian	Black	Other	Not given	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	29%	*	*	*	*	7%	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	79%	*	*	*	*	87%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	68%	*	*	*	*	64%	68%

YOUR OVERALL NHS CARE		Ethnicity					
	White	Mixed	Asian	Black	Other	Not given	All
Q56. The whole care team worked well together	90%	*	*	*	*	98%	91%
Q57. Administration of care was very good or good	88%	*	*	*	*	84%	88%
Q58. Cancer research opportunities were discussed with patient	44%	*	*	*	*	27%	42%
Q59. Patient's average rating of care scored from very poor to very good	9.0	*	*	*	*	8.8	9.0

IMD quintile tables

SUPPORT FROM YOUR GP PRACTICE	IMD quintile						
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	74%	76%	75%	85%	89%	*	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	46%	63%	63%	69%	70%	*	63%

DIAGNOSTIC TESTS	IMD quintile								
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All		
Q5. Patient received all the information needed about the diagnostic test in advance	90%	91%	93%	93%	96%	*	92%		
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	74%	87%	87%	85%	83%	*	85%		
Q7. Patient felt the length of time waiting for diagnostic test results was about right	71%	78%	76%	81%	70%	*	76%		
Q8. Diagnostic test results were explained in a way the patient could completely understand	74%	81%	79%	81%	83%	*	80%		
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	89%	98%	95%	95%	95%	*	95%		

FINDING OUT THAT YOU HAD CANCER			IMD quintile				
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	81%	83%	80%	80%	80%	*	81%
Q13. Patient was definitely told sensitively that they had cancer	70%	75%	78%	75%	76%	*	75%
Q14. Cancer diagnosis explained in a way the patient could completely understand	71%	79%	76%	80%	79%	*	77%
Q15. Patient was definitely told about their diagnosis in an appropriate place	78%	87%	83%	87%	86%	*	85%
Q16. Patient was told they could go back later for more information about their diagnosis	84%	88%	89%	90%	85%	*	88%

SUPPORT FROM A MAIN CONTACT PERSO	N		IM	ID quintile			
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q17. Patient had a main point of contact within the care team	86%	94%	92%	92%	95%	*	92%
Q18. Patient found it very or quite easy to contact their main contact person	87%	81%	86%	88%	88%	*	86%
Q19. Patient found advice from main contact person was very or quite helpful	94%	97%	96%	96%	96%	*	96%

IMD quintile tables

DECIDING ON THE BEST TREATMENT			IIV	1D quintile			
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q20. Treatment options were explained in a way the patient could completely understand	73%	86%	88%	84%	86%	*	84%
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	74%	82%	84%	77%	89%	*	81%
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	73%	87%	84%	80%	91%	*	84%
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	62%	62%	54%	52%	56%	*	57%

CARE PLANNING	IMD quintile							
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All	
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	74%	74%	72%	73%	77%	*	73%	
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	90%	91%	95%	93%	95%	*	93%	
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	98%	99%	100%	99%	100%	*	99%	

SUPPORT FROM HOSPITAL STAFF	IMD quintile						
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q27. Staff provided the patient with relevant information on available support	92%	96%	94%	93%	96%	*	94%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	79%	80%	82%	78%	82%	*	80%
Q29. Patient was offered information about how to get financial help or benefits	55%	60%	69%	70%	74%	*	65%

HOSPITAL CARE			IIV	1D quintile			
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	76%	78%	70%	79%	74%	*	76%
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	56%	74%	61%	67%	71%	*	67%
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	71%	74%	69%	75%	74%	*	73%
Q34. Patient was always able to get help from ward staff when needed	71%	72%	69%	75%	72%	*	72%
Q35. Patient was always able to discuss worries and fears with hospital staff	65%	67%	69%	63%	67%	*	66%
Q36. Hospital staff always did everything they could to help the patient control pain	82%	83%	84%	85%	88%	*	84%
Q37. Patient was always treated with respect and dignity while in hospital	86%	87%	87%	87%	96%	*	88%
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	78%	87%	81%	80%	76%	*	82%
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	82%	86%	86%	83%	78%	*	84%

IMD quintile tables

YOUR TREATMENT			IIV	1D quintile			
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	86%	90%	90%	87%	89%	*	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	91%	87%	85%	83%	83%	*	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	84%	81%	90%	85%	86%	*	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	73%	85%	84%	74%	60%	*	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	95%	95%	79%	89%	89%	*	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	84%	80%	82%	88%	79%	*	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	88%	83%	83%	77%	81%	*	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	84%	75%	80%	76%	75%	*	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	73%	87%	73%	71%	60%	*	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	86%	85%	82%	78%	81%	*	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	81%	78%	77%	80%	77%	*	78%

IMMEDIATE AND LONG TERM SIDE EFFEC	TS		IM	1D quintile			
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	69%	77%	79%	74%	71%	*	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	75%	73%	70%	68%	76%	*	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	81%	88%	91%	87%	89%	*	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	56%	63%	68%	65%	54%	*	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	50%	54%	63%	61%	48%	*	56%

SUPPORT WHILE AT HOME	IMD quintile							
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All	
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	65%	68%	65%	65%	59%	*	65%	
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	59%	44%	61%	55%	43%	*	52%	

CARE FROM YOUR GP PRACTICE		IMD quintile					
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q51. Patient definitely received the right amount of support from their GP practice during treatment	42%	42%	53%	50%	43%	*	47%
Q52. Patient has had a review of cancer care by GP practice	29%	17%	21%	17%	20%	*	20%

IMD quintile tables

LIVING WITH AND BEYOND CANCER		IMD quintile					
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	18%	16%	26%	52%	9%	*	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	77%	79%	87%	77%	75%	*	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	58%	67%	62%	76%	75%	*	68%

YOUR OVERALL NHS CARE		IMD quintile					
	1 (most deprived)	2	3	4	5 (least deprived)	Non- England	All
Q56. The whole care team worked well together	91%	89%	94%	89%	91%	*	91%
Q57. Administration of care was very good or good	88%	85%	90%	89%	85%	*	88%
Q58. Cancer research opportunities were discussed with patient	37%	39%	46%	46%	41%	*	42%
Q59. Patient's average rating of care scored from very poor to very good	8.9	8.9	9.0	8.9	9.0	*	9.0

SUPPORT FROM YOUR GP PRACTICE		Long-term con		
	Yes	No	Not given	All
Q2. Patient only spoke to primary care professional once or twice before cancer diagnosis	77%	84%	75%	79%
Q3. Referral for diagnosis was explained in a way the patient could completely understand	59%	70%	68%	63%

DIAGNOSTIC TESTS		Long-term con-	dition status	
	Yes	No	Not given	All
Q5. Patient received all the information needed about the diagnostic test in advance	92%	93%	87%	92%
Q6. Diagnostic test staff appeared to completely have all the information they needed about the patient	83%	90%	78%	85%
Q7. Patient felt the length of time waiting for diagnostic test results was about right	77%	75%	73%	76%
Q8. Diagnostic test results were explained in a way the patient could completely understand	80%	83%	66%	80%
Q9. Enough privacy was always given to the patient when receiving diagnostic test results	95%	96%	93%	95%

FINDING OUT THAT YOU HAD CANCER		Long-term con		
	Yes	No	Not given	All
Q12. Patient was told they could have a family member, carer or friend with them when told diagnosis	82%	80%	83%	81%
Q13. Patient was definitely told sensitively that they had cancer	76%	74%	70%	75%
Q14. Cancer diagnosis explained in a way the patient could completely understand	77%	78%	74%	77%
Q15. Patient was definitely told about their diagnosis in appropriate place	85%	85%	78%	85%
Q16. Patient was told they could go back later for more information about their diagnosis	88%	90%	76%	88%

SUPPORT FROM A MAIN CONTACT PERSO	Long-term condition status			
	Yes	No	Not given	All
Q17. Patient had a main point of contact within the care team	93%	90%	90%	92%
Q18. Patient found it very or quite easy to contact their main contact person	85%	86%	89%	86%
Q19. Patient found advice from main contact person was very or quite helpful	96%	97%	94%	96%

DECIDING ON THE BEST TREATMENT	Long-term condition status					
	Yes No Not given					
Q20. Treatment options were explained in a way the patient could completely understand	84%	86%	82%	84%		
Q21. Patient was definitely involved as much as they wanted to be in decisions about their treatment	81%	82%	80%	81%		
Q22. Family and/or carers were definitely involved as much as the patient wanted them to be in decisions about treatment options	84%	85%	76%	84%		
Q23. Patient could get further advice from a different healthcare professional before making decisions about their treatment options	55%	61%	59%	57%		

CARE PLANNING	Long-term condition status					
	Yes No Not given All					
Q24. Patient was definitely able to have a discussion about their needs or concerns prior to treatment	74%	73%	71%	73%		
Q25. A member of their care team helped the patient create a care plan to address any needs or concerns	92%	95%	88%	93%		
Q26. Care team reviewed the patient's care plan with them to ensure it was up to date	100%	98%	100%	99%		

SUPPORT FROM HOSPITAL STAFF	Long-term condition status			
	Yes	No	Not given	All
Q27. Staff provided the patient with relevant information on available support	94%	95%	96%	94%
Q28. Patient definitely got the right level of support for their overall health and well being from hospital staff	80%	81%	77%	80%
Q29. Patient was offered information about how to get financial help or benefits	62%	75%	53%	65%

HOSPITAL CARE	Long-term condition status				
	Yes	No	Not given	All	
Q31. Patient had confidence and trust in all of the team looking after them during their stay in hospital	74%	80%	75%	76%	
Q32. Patient's family, or someone close, was definitely able to talk to a member of the team looking after the patient in hospital	64%	73%	63%	67%	
Q33. Patient was always involved in decisions about their care and treatment whilst in hospital	68%	80%	76%	73%	
Q34. Patient was always able to get help from ward staff when needed	70%	75%	76%	72%	
Q35. Patient was always able to discuss worries and fears with hospital staff	63%	70%	77%	66%	
Q36. Hospital staff always did everything they could to help the patient control pain	83%	88%	81%	84%	
Q37. Patient was always treated with respect and dignity while in hospital	87%	88%	92%	88%	
Q38. Patient received easily understandable information about what they should or should not do after leaving hospital	78%	89%	83%	82%	
Q39. Patient was always able to discuss worries and fears with hospital staff while being treated as an outpatient or day case	82%	87%	84%	84%	

YOUR TREATMENT		Long-term co	ndition status	
	Yes	No	Not given	All
Q41_1. Beforehand patient completely had enough understandable information about surgery	88%	90%	89%	89%
Q41_2. Beforehand patient completely had enough understandable information about chemotherapy	85%	86%	89%	86%
Q41_3. Beforehand patient completely had enough understandable information about radiotherapy	83%	87%	94%	85%
Q41_4. Beforehand patient completely had enough understandable information about hormone therapy	71%	88%	*	78%
Q41_5. Beforehand patient completely had enough understandable information about immunotherapy	88%	88%	*	89%
Q42_1. Patient completely had enough understandable information about their response to surgery	81%	87%	81%	83%
Q42_2. Patient completely had enough understandable information about their response to chemotherapy	81%	84%	85%	82%
Q42_3. Patient completely had enough understandable information about their response to radiotherapy	72%	85%	87%	78%
Q42_4. Patient completely had enough understandable information about their response to hormone therapy	72%	80%	*	75%
Q42_5. Patient completely had enough understandable information about their response to immunotherapy	80%	84%	*	82%
Q43. Patient felt the length of waiting time at clinic and day unit for cancer treatment was about right	78%	80%	77%	78%

IMMEDIATE AND LONG TERM SIDE EFFECTS		Long-term con	dition status	
	Yes	No	Not given	All
Q44. Possible side effects from treatment were definitely explained in a way the patient could understand	73%	79%	75%	75%
Q45. Patient was always offered practical advice on dealing with any immediate side effects from treatment	70%	76%	65%	72%
Q46. Patient was given information that they could access about support in dealing with immediate side effects from treatment	86%	92%	84%	88%
Q47. Patient felt possible long-term side effects were definitely explained in a way they could understand in advance of their treatment	60%	70%	58%	63%
Q48. Patient was definitely able to discuss options for managing the impact of any long-term side effects	53%	64%	51%	56%

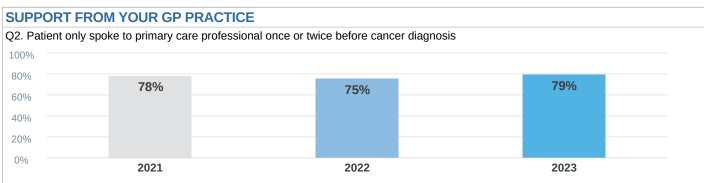
SUPPORT WHILE AT HOME	Long-term condition status				
	Yes No Not given				
Q49. Care team gave family, or someone close, all the information needed to help care for the patient at home	62%	74%	57%	65%	
Q50. During treatment, the patient definitely got enough care and support at home from community or voluntary services	51%	64%	39%	52%	

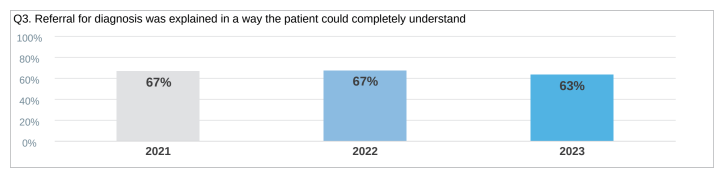
CARE FROM YOUR GP PRACTICE	Long-term condition status				
	Yes No Not given All				
Q51. Patient definitely received the right amount of support from their GP practice during treatment	43%	54%	49%	47%	
Q52. Patient has had a review of cancer care by GP practice	20%	18%	27%	20%	

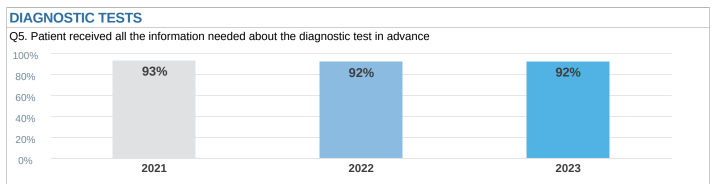
LIVING WITH AND BEYOND CANCER	Long-term condition status			
	Yes	No	Not given	All
Q53. After treatment, the patient definitely could get enough emotional support at home from community or voluntary services	28%	33%	6%	26%
Q54. The right amount of information and support was offered to the patient between final treatment and the follow up appointment	77%	86%	75%	79%
Q55. Patient was given enough information about the possibility and signs of cancer coming back or spreading	64%	75%	64%	68%

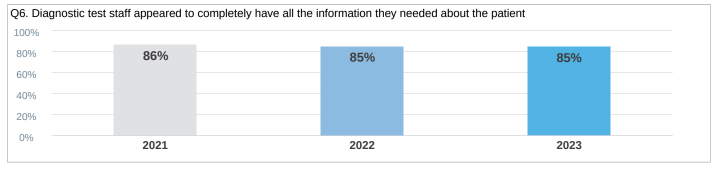
YOUR OVERALL NHS CARE	Long-term condition status				
	Yes No Not given A				
Q56. The whole care team worked well together	90%	91%	95%	91%	
Q57. Administration of care was very good or good	88%	89%	80%	88%	
Q58. Cancer research opportunities were discussed with patient	39%	52%	35%	42%	
Q59. Patient's average rating of care scored from very poor to very good	8.9	9.1	8.7	9.0	

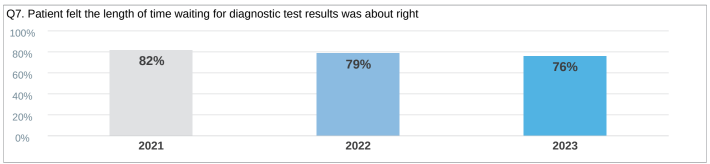


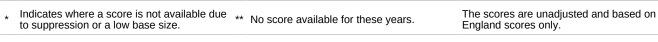


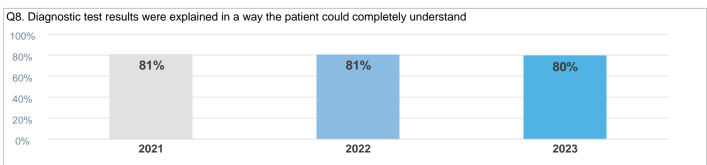


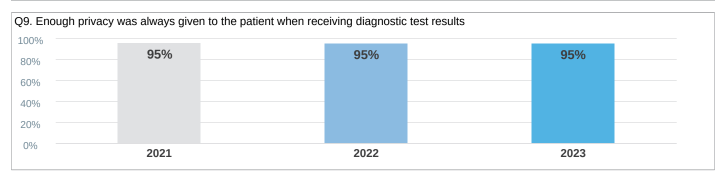


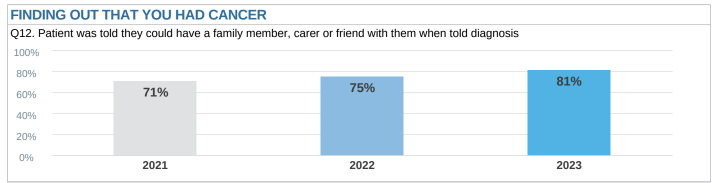


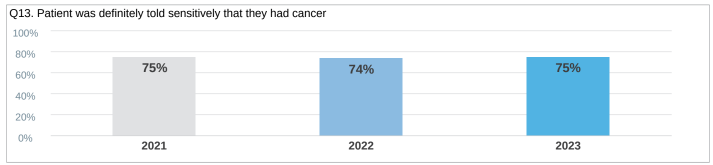


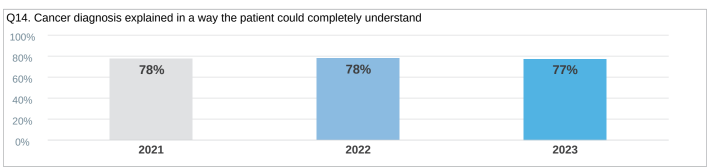


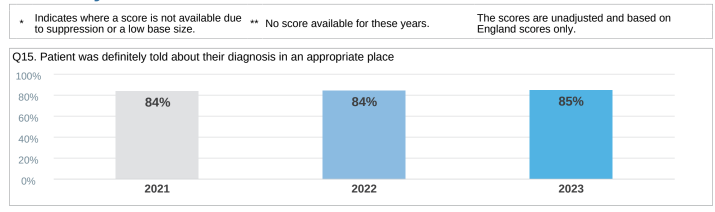


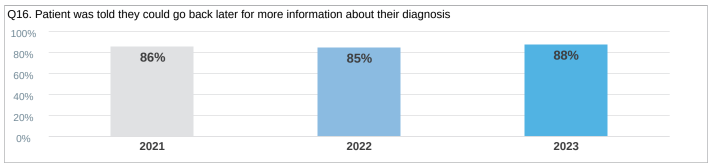


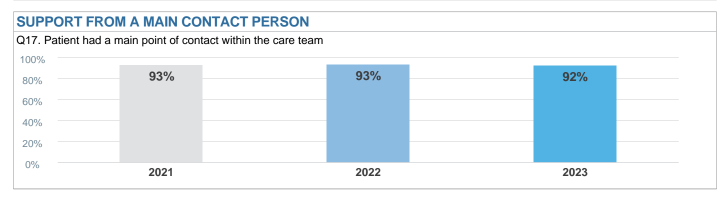


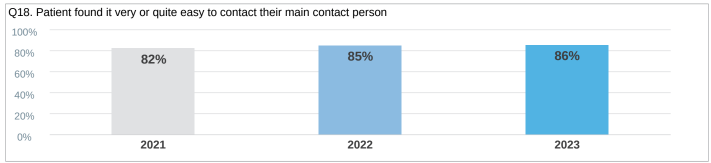


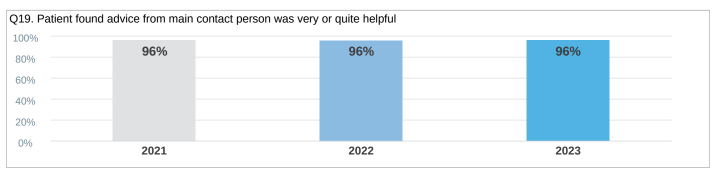




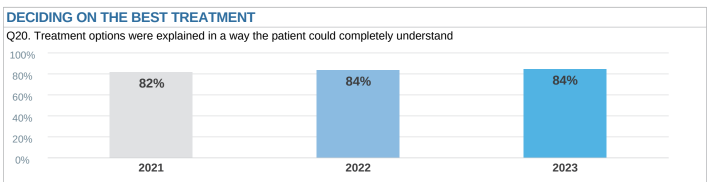


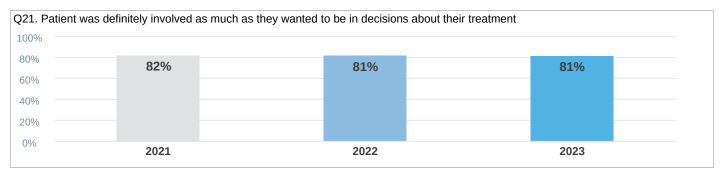


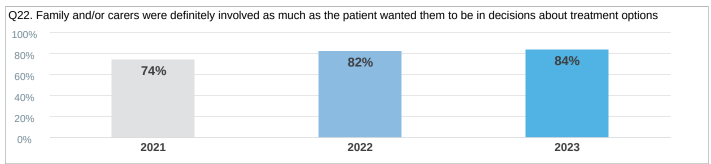


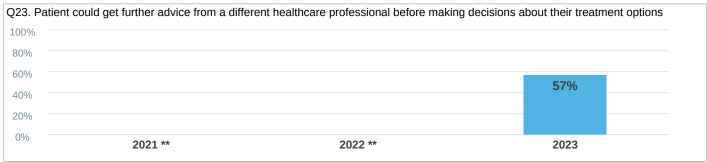


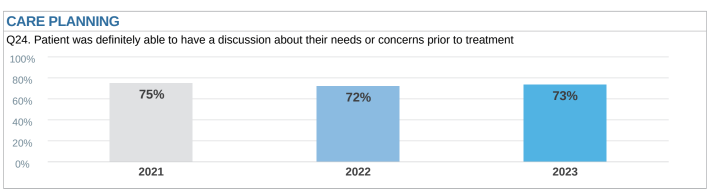


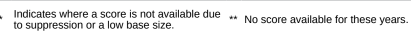




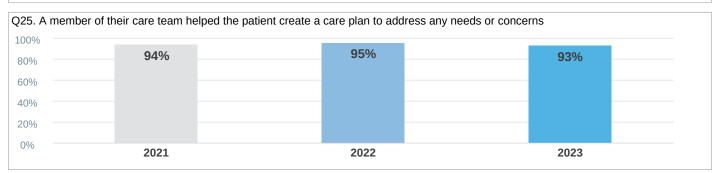


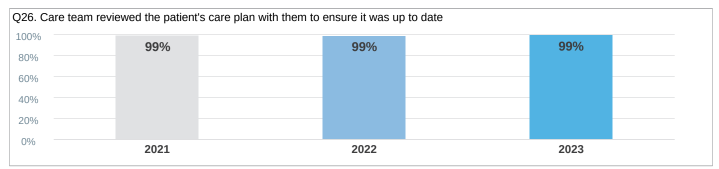


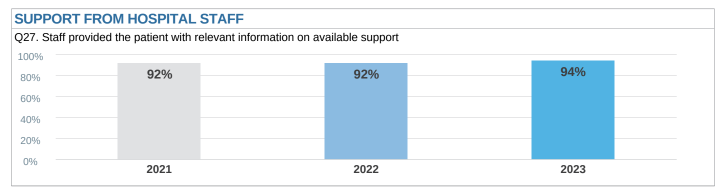


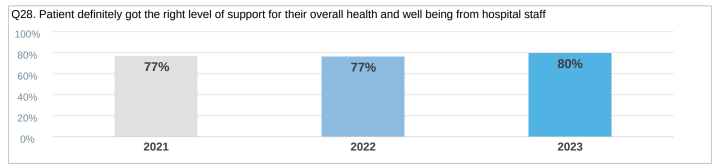


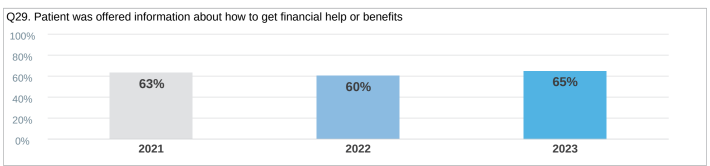
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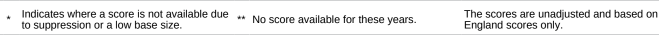


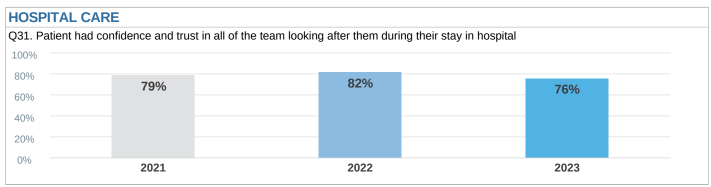


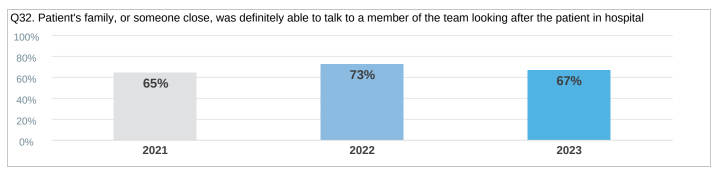


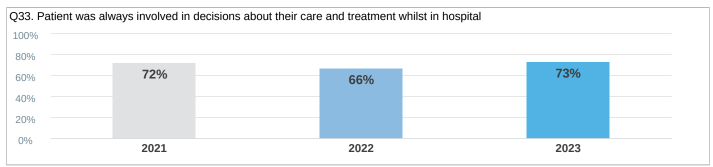


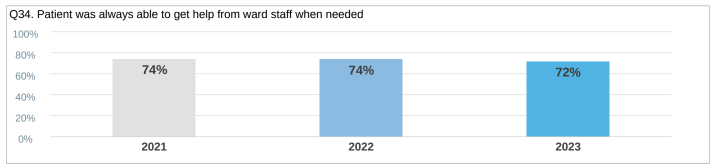


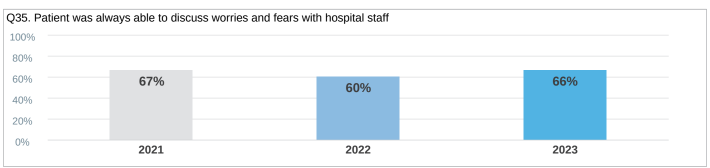


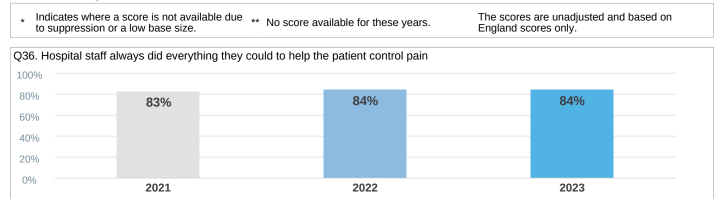


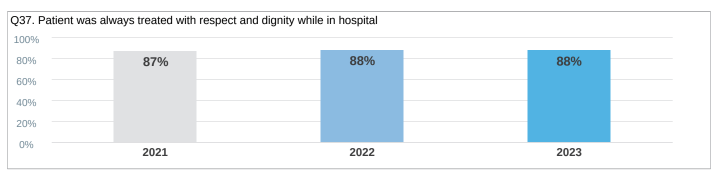


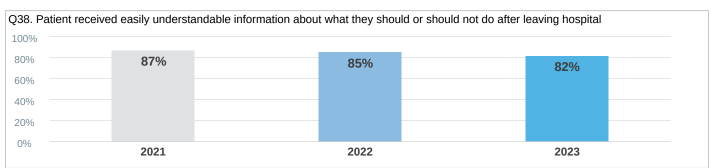


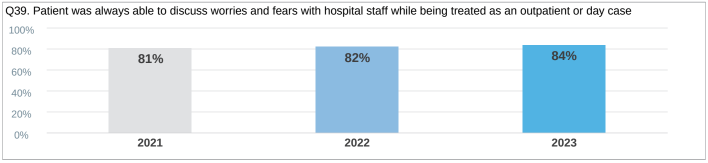


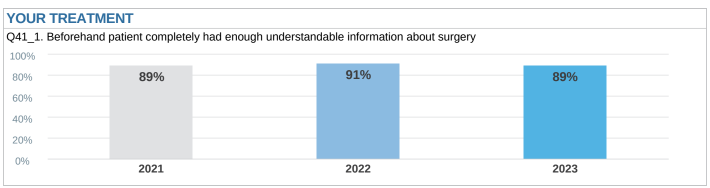


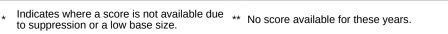




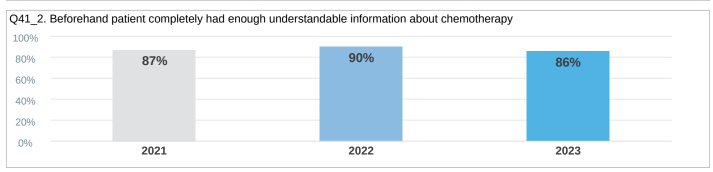


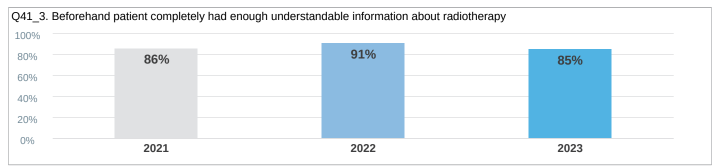


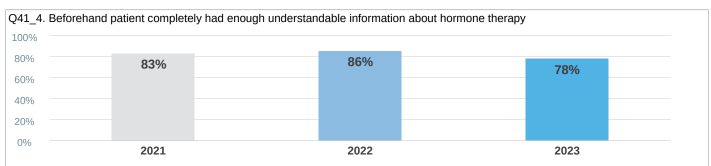


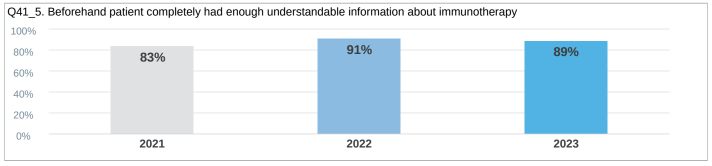


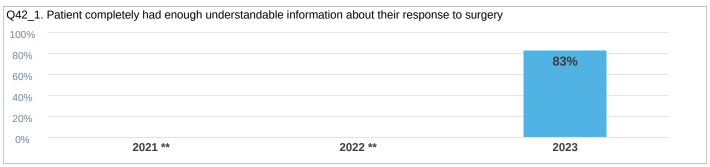
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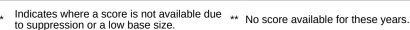
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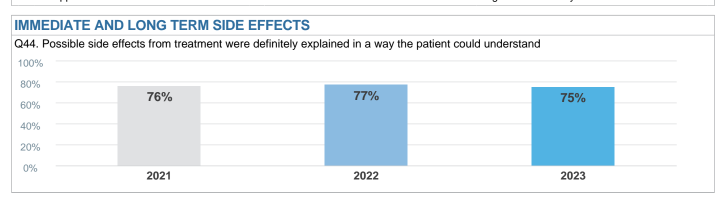


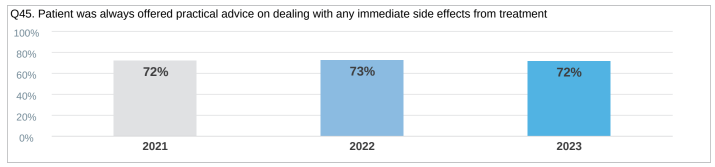
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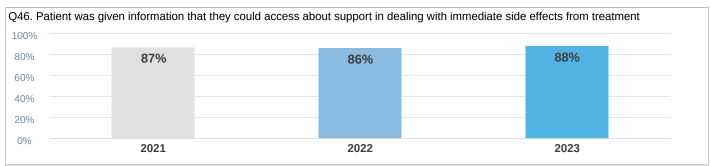
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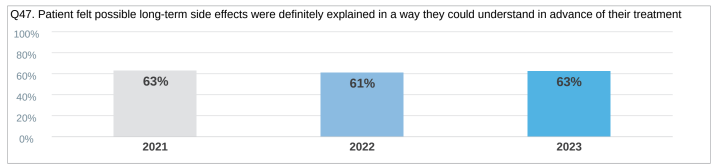


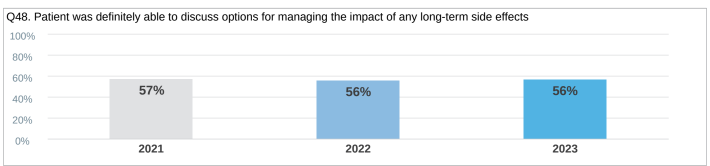
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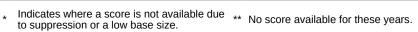




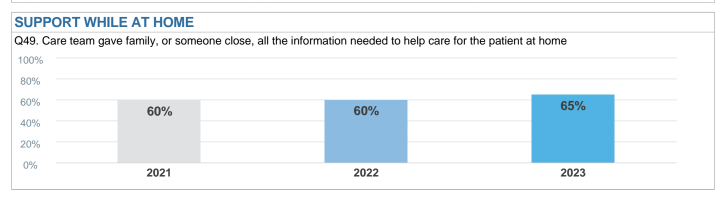


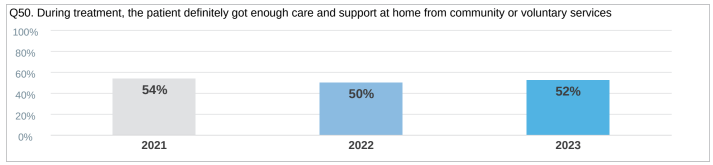


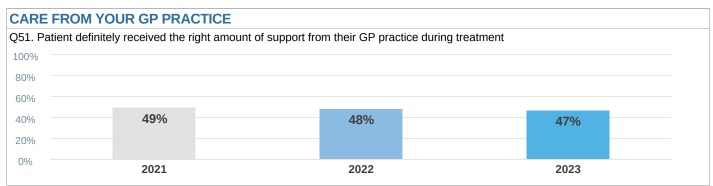


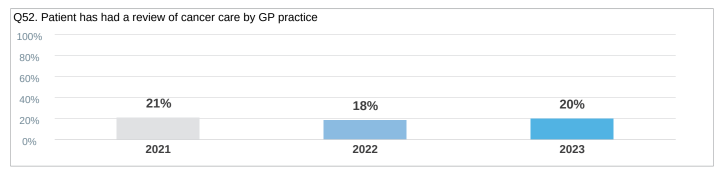


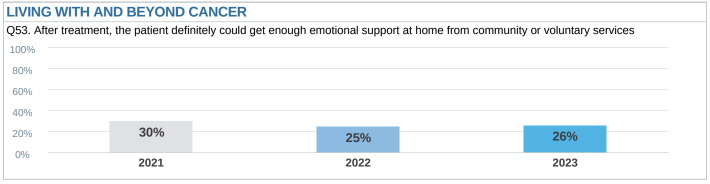
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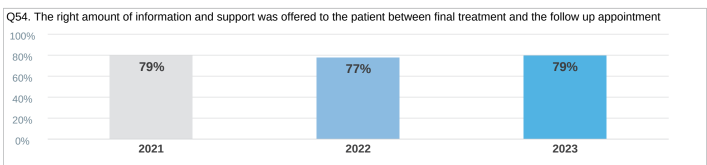


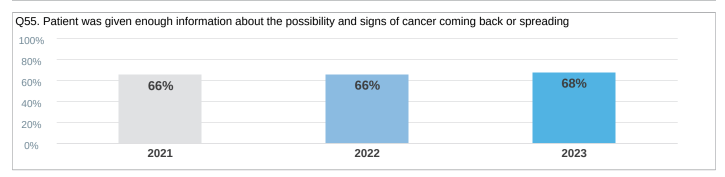


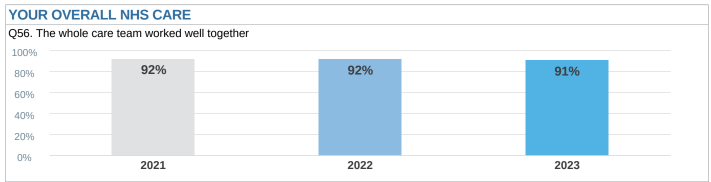


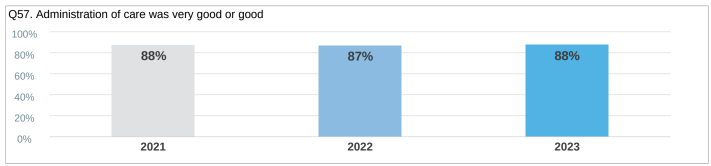


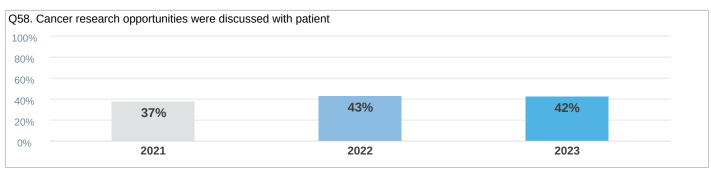












Cancer Patient Experience Survey 2023 University Hospitals Plymouth NHS Trust

